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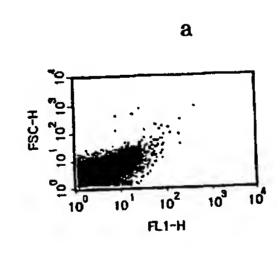
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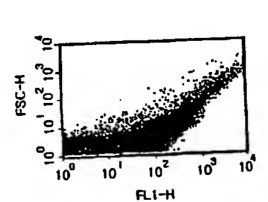
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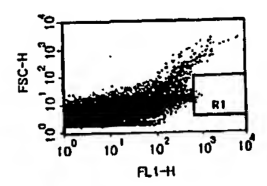


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(57) Abstract: The present invention relates to a method for display of proteinson spore surface and a method for improving protein with rapidity using the same, which comprises the steps of (i) preparing a vector for spore surface display comprising a gene construct containing a gene encoding spore coat protein and a gene encoding a protein of interest, wherein, when expressed, the gene construct expresses a fusion protein between the spore coat protein and the protein of interest, (ii) transforming a host cell with the vector for spore surface display; (iii) displaying the protein of interest on a surface of a spore of the host cell; and (iv) recovering the spore displaying on its surface the protein of interest.



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METHOD FOR EXPRESSION OF PROTEINS ON SPORE SURFACE

FIELD OF THE INVENTION

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The present invention relates to a method for display of proteins on spore surface, in particular to a method for surface display using spore coat proteins as surface display motif and a high throughput method for improving protein.

DESCRIPTION OF THE RELATED ART

The technology of surface display in which organism displays on its surface the desired proteinaceous substance such as peptide and polypeptide has wider application fields depending on the types of protein displayed or host organism (Georgiou et al., 1993, 1997; Fischetti et al., 1993; and Schreuder et al., 1996). The gene of protein to be displayed is contained in host organism and thus the host can be selectively screened using the characteristics of the protein displayed, thereby obtaining the desired gene from the selected host with easiness. Therefore, such surface display technology can guarantee a powerful tool on molecular evolution of protein (see WO 9849286; and U.S. Pat. No. 5,837,500).

High-Throughput Screening

For instance, phage displaying on its surface antibody

having desired binding affinity is bound to immobilized antigen and then eluted, followed by propagating the eluted phage, thereby yielding the gene coding for target antibody from phage (U.S. Pat. No. 5,837,500). The bio panning method described above can provide a tool to select target antibody by surface displaying antibody library on phage surface in large amount and comprises the consecutive steps as follows:

(1) constructing library; (2) surface displaying the library; (3) binding to immobilized antigen; (4) eluting the bound phage; finally (5) propagating selected clones.

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The technology of phage surface display has been found to be useful in obtaining the desired monoclonal variant form enormous library (e.g., 106-109 variants) and thus applied to the field of high-throughput screening of antibody. Antibody has been used in various fields such as therapy, diagnosis, etc. and thus its demand has been increased. In this context, there has been a need for novel antibody to have binding affinity to new substance or catalyze biochemical reaction. The hybridoma technology to produce monoclonal antibody has been conventionally used so as to satisfy the need. However, the conventional method needs high expenditure and long time for performance whereas the yield of antibody is very low. In addition to this, to screen novel antibody, more than 10^{10} antibody libraries is generally used, as a result, the hybridoma technology has

been thought to be inadequate in finding antibody exhibiting new binding property.

Many researches has focused on novel methods which is easier and more effective that the bio panning method described above and then developed novel technologies performed in such a manner that libraries are displayed on surface of bacteria or yeast and then cells displaying target protein is sorted with flow cytometry in a highthroughput manner. According to the technology, antigen labeled with fluorescent dye is bound to surface-displaying cell and the antibody having the desired binding affinity is isolated with flow cytometry capable of analyzing more than 108 cells a hour. Francisco, et al., have demonstrated the usefulness of microbial display technology by revealing that surface-displayed monoclonal antibody could be concentrated with flow cytometry at rate of more than 105, finally more than 79% have been proved to be the desired cells (Daugherty et al., 1998).

Live Vaccine

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The surface display technology mentioned above can display antigen or fragment thereof and hence provide a delivery system for recombinant live vaccine. Up to now, attenuated pathogens or viruses have been predominantly employed as vaccine. Particularly, the bacteria have been

found to express antigen intracellularly or extracellularly or on its cell membrane, thereby delivering antigen to host cell. The surface-displayed live vaccine induces a potential immune reaction and expresses continuously antigen during propagation in host cell; therefore, it has been highlighted as novel delivery system for vaccine. In particular, pathogen-derived antigenic epitope displayed on surface of nonpathogenic *E. coli* or Salmonella is administered orally in viable form and then exhibits to induce immune reaction in more continuous and powerful manner (Georgiou et al., 1997; and Lee et al., 2000).

Whole Cell Bioconversion

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Whole cell as biocatalyst displaying on its surface enzyme capable of catalyzing chemical reaction can avoid expression, isolation direct and for necessities stabilization of enzyme. In case of expressing enzyme in cell for bioconversion, the cell is compelled to recovery (e.g., toluene) treatment ensure chemical to and impermeability of substrate. In addition, the lasting use inactive or gives a problem on renders the enzyme transference of substrate and product, thus dropping the productivity of overall process.

The above-mentioned shortcomings can be removed using enzyme displayed on cell surface (Jung et al, 1998a: 1998b).

With whole cell displaying on its surface phosphodiesterase, organophosphorous-typed parathion and paraoxon with higher toxicity can be degraded, which is a typical example representing the applicability of cells displaying enzyme to environmental purification process (Richins et al., 1997).

Antipeptide Antibody

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Martineau et al. have reported a highly simple method for production of antipeptide antibody using surface display technology of *E. coli* (Martineau et al., 1991). As described, the desired peptide is displayed on the protruding region of MalE and outer membrane protein, LamB and then whole cell or fragmented cell is administered to animal so as to generate antipeptide antibody. The method makes it possible to produce antibody with avoiding chemical synthesis of peptide and its linkage to carrier protein.

Whole Cell Absorber

To immobilize antibody or polypeptide on suitable carrier, which is useful in absorption chromatography, several subsequent steps must be performed, for example, protein production by fermentation, isolation of protein in pure form, and immobilization on a carrier. Generally, it is difficult to prepare the bioabsorber.

As absorber, a whole cell displaying absorption protein

has been developed. The whole cell absorber known mostly is Staphylococcus aureus displaying on its surface protein A naturally, which has a high binding affinity to Fc domain of mammalian antibody. Currently, novel method has been proposed to remove and recover heavy metals, which employs metallothionein or metal-absorption protein displayed on microbial cell surface in large amount (Sousa et al., 1996, 1998; and Samuelson et al., 2000). The method is more effective in removing and recovering heavy metals from contamination source in comparison with the conventional method using metal-absorption microbes.

As understood based on the matters described above, in order to display foreign protein on cell surface, a suitable surface protein and foreign protein must be linked each other in gene level to express fusion protein, and the fusion protein should pass stably across inner membrane of cell to be attached to cell surface. Preferably, the surface protein having the following characteristics is recommended as surface display motif: 1) existence of secretory signal enabling passage across inner membrane of cell, 2) existence of target signal enabling stable attachment to cell surface, 3) high expression level on cell surface, and 4) stable expression regardless of protein size (Georgiou et al., 1993).

Therefore, the surface display motif or novel recombinant protein, which meets the requirements described above, should be selected or prepared to develop novel surface display system overcoming disadvantages of the known systems. In addition, the selection of a suitable host cell to display is very pivotal.

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Up to date, the developed surface display systems are as follows: phage surface display system (Chiswell and McCarferty, 1992), bacterial surface display system (Georgiou et al., 1993; Little et al., 1993; and Georgiou et al., 1997), surface display system of Gram negative bacteria (Francisco et al., 1992; Fuchs et al., 1991; Klauser et al., 1990, 1992; and Hedegaard et al., 1989), surface display system of Gram positive bacteria (Samuelson et al., 1995; Palva et al., 1994; and Sleytr and Sara, 1997), and surface display system of yeast (Ferguson, 1988; and Schreuder et al., 1996).

In the developed phage display system, the concentration of the desired clone from phage library has been found to be difficult and the antibody selected from phage library displaying has usually exhibited very low expression rate. According to a surface display system of Gram negative bacteria, the incorporation of foreign polypeptide into surface structure results in not only its steric limitation which makes it impossible to have stable membrane protein

(Charbit et al., 1987; and Agterberg et al., 1990) but also drop of the stability of cell outer membrane and its viability. In addition, in surface display system of yeast, because the vector used has usually shown a low rate of transformation, which is unfavorable to surface display of library.

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The surface display systems developed have been cooperatively used each other. For example, to screen antibody variant with enhanced binding affinity, a primary screening is performed using phage surface display system and additionally, the secondary screening is carried out using cell surface display system (Georgiou, 2000). However, the phage display technology is encountered to difficulty in concentration of the desired clones from phage library. The reason is that the antibody displayed on phage surface does not show the elution pattern depending exactly on its binding affinity, which is ascribed to avidity of antibody displayed on phage surface. Therefore, there remains a need of novel methods ensuring screening the desired antibody from antibody library.

E. coli as display host, which has been intensively studied, uses generally cell outer membrane protein as surface display motif. However, the over-expression of cell outer membrane protein fused to foreign protein is likely to

bring about structural instability of cell outer membrane, consequently, diving the viability of host cell (Georgiou et al., 1996). To be from the shortcomings, ice-nucleation protein with no effect on viability has been used as display motif, and has been applied to bioconversion process, surface display of enzyme library and screening enzyme variants (Jung et al., 1998a, 1998b; and Kim et al., 1998, 1999, 2000).

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The size of library displayed on surface depends on the transformation efficiency of host cell with vector; thus *E. coli* as host has an advantage in view of the size of library to be displayed. Gram positive bacteria as host are relatively rigid and permit stable display of the desired protein; however, transformation efficiency is exhibited low, which results in smaller size of library than *E. coli*.

The host organisms having been developed are likely to be sensitive to a variety of physiochemical treatments, which makes it impossible to select proteins displayed on surface by virtue of direct physiochemical treatment. For example, in screening a variant of antibody with enhanced binding affinity, abrupt change of pH or adjustment of the concentration of base is generally performed to elute the variant, which are found to decrease the viability of phage or bacteria in medium.

In addition, the host organisms used conventionally have

a complicated and weak structure of cell surface, which drops adaptability to extreme environment such as high temperature and high pressure. To employ *E. coli* displaying on its surface enzyme in bioconversion reaction, the cells must have represent stability in bioconversion system. In this context, the surface of *E. coli* displaying on its surface enzyme is generally subject to immobilization, which does not lead to satisfying results (Freeman et al., 1998).

display known surface above, the described As based on applying fields, used have technologies, bacteriophage, Gram negative or positive bacterium, yeast, cilium or mammalian cell as host organism and surface proteins of each organism as surface display motif. However, in the surface display methods having been developed, the host organism does not have resistance to chemicals and physiochemical change such as pH change, and displaying protein on its surface in excess leads to disadvantages in cell surface, finally reducing the viability of host cell largely (Georgiou et al., 1996).

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DETAILED DESCRIPTION OF THIS INVENTION

Under such situation, the present inventors have made intensive studies to be from the shortcoming of conventional display methods, and as a result, we have developed novel display system using a spore as host and a coat protein as

motif of surface display. Surprisingly, the developed display system has been found to have excellent stability to a variety of physiochemical stresses in surrounding environment and have much broader applicability.

Accordingly, it is an object of this invention to provide a method for displaying a protein of interest on spore surface using a system for spore surface display.

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It is another object of this invention to provide a method for improving a protein of interest using a system for spore surface display.

It is still another object of this invention to provide a method for bioconversion using a system for spore surface display.

It is further object of this invention to provide a method for preparing protein microarray using a system for spore surface display.

It is still further object of this invention to provide a method producing an antibody to antigen in vertebrates using a system for spore surface display.

It is another object of this invention to provide a method for preparing a whole cell absorber using a system for spore surface display.

It is still another object of this invention to provide a microbial transformant for spore surface display of a protein of interest.

It is further object of this invention to provide a spore for spore surface display of a protein of interest.

It is still further object of this invention to provide a vector for spore surface display.

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The principle of the present invention lies in the employment of microbial spore as host for surface display and spore coat protein as surface display motif. The present inventors have been compelled to select a system for spore surface display since the the spore has a following advantages (Driks, 1999): 1) a higher heat stability, 2) a significant stability to radioactivity, 3) a stability to toxins, 4) a higher stability to acid and base, 5) a significant stability to lysozyme, 6) a resistance to dryness, 7) a higher stability to organic solvents, 8) a fusion protein between a surface display motif and a protein of interest is displayed on spore surface immediately after expression without secretion in host cell, 9) no metabolic activity, and 10) shorter time for obtaining spore, e.g. within several hours.

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In particular, the spore coat proteins used in this invention circumvent a necessity for passage across cell membrane, so that they do not need secretion signal and target signal which are prerequisites of surface display motif, thereby ensuring a surface display of protein such as

 β -galactosidase, in orderly fashion, which is difficult to pass across cell membrane.

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U.S. Pat. No. 5,766,914 discloses a method of producing and purifying enzymes using fusion protein between cotC or cotD among spore coat proteins of Bacillus subtilis and lacZ as reporter. However, as disclosed, a purification method demonstrating surface display of protein is not recognized to isolate spores specifically. Furthermore, the activity of enzyme expressed has been very low and the display of enzyme on spore surface has never demonstrated by means of reliable methods such biochemical, physical and immunological methods. In addition to this, the inner coat protein, cotD is enclosed by outer coat protein of 70-200 nm thickness, which makes it difficult to be exposed to spore surface. In case of fusion protein expression using outer coat protein, cotC, the activity of enzyme is increased by four-fold in comparison with that of cotD; however, the activity, 0.02 U, is considered negligible, in particular, in consideration of industrial scale. Therefore, the matter disclosed in the document above cannot be considered to use and recognize a system for spore surface display. In other words, the patent document cannot be recognized to describe a system for spore surface display. U.S. Pat. Nos. 5,837,500 and 5,800,821 also indicate cotC and cotD as a preferable surface display motif,

and therefore the patent documents cannot be recognized to describe a system for spore surface display because of the reasons mentioned above.

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Furthermore, according to the purification method of spore proposed in U.S. Pat. No. 5,766,914, half of the purified resultant has been observed under microscope to have the complex forms between cells harboring spores and cell-lysis matters bound to spores (see Fig. 1; cells with blackish color and long side are those not forming spore and spores is observed to be white and circular), which has been demonstrated by the present inventors. The facts hereinabove reveals possibility to bring about the false results by measuring of the activity of reporter enzyme or analyzing of reporter enzyme with flow cytometry in vegetative cells rather than on spore surface. In contrast, the renografin gradient centrifugation as demonstrated in Examples below allows for the perfect purification of spores (see Fig. 2), thereby measuring the activity of enzyme displayed on spore surface solely.

Observations on lower enzyme activity in several documents including the patents above are likely to be resulted from the following reasons. First, it is suggested that the expression level of coat protein itself is low. The maximum expression levels of CotC and CotD are 40 and 147 Miller Units, respectively, which is considered to be

largely low, in particular, in consideration of CotE of 6021 Miller Units (Zheng L and Losick R., J. Mol. Biol. 212:645-660(1990)). Furthermore, it is notable that the amount of enzyme displayed on spore surface has not been reported. Secondly, it is possible that the protein displayed on spore surface is cleaved by protease in host cell. Such suggestion is made based on the fact that at spore-forming stage of Bacillus subtilis a variety of proteases are expressed and reconstitution for spore formation is occurred. The suggestion can be demonstrated in Examples below in which a variant lack protease exhibits a much higher enzyme activity displayed on spore surface (see Fig. 7).

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Using gene of GFP (green fluorescence protein) as reporter linked to cotE and spoIVA, the studies on gene expression and localization of the expressed protein in spore has been attempted (Webb et al., 1995; Lewis et al., 1996). The publications disclose that the fusion protein expressed is found in spore by means of observation under fluorescence microscope using fluorescence of GFP; however, they never describe if the fusion protein is displayed and linked on spore surface.

As another example of spore surface display using coat protein, U.S. Pat. No. 5,800,821 discloses a spore as delivery system of antigen. However, the publication does not disclose that the antigen expressed is displayed on

spore surface and the spore containing antigen administered can induce immunization reaction in host.

The present inventors have recognized the shortcomings of the conventional arts described above and developed an efficient and optimized system for spore surface display, which have been confirmed by enzymological, immunological and physiochemical methods using various spore coat proteins.

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In one aspect of this invention, there is provided a method for displaying a protein of interest on spore surface, which comprises the steps of: (i) preparing a vector for spore surface display comprising a gene construct containing a gene encoding spore coat protein and a gene encoding a protein of interest, wherein, when expressed, the gene construct expresses a fusion protein between the spore coat protein and the protein of interest; (ii) transforming a host cell with the vector for spore surface display; (iii) displaying the protein of interest on a surface of a spore of the host cell; and (iv) recovering the spore displaying on its surface the protein of interest.

In another aspect of this invention, there is provided a method for improving a protein of interest, which comprises the steps of: (i) constructing a gene library of the protein of interest; (ii) preparing a vector by linking the gene

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library to a gene encoding spore coat protein; (iii) transforming a spore-forming host cell with the vector; (iv) forming a spore in the transformed host cell and displaying the protein of interest on a surface of the spore; (v) recovering the spore displaying on its surface the protein of interest; and (vi) screening the spore displaying a variant of the protein of interest having a desired property.

In still another aspect of this invention, there is provided a method for improving a protein of interest using a resistance property of spore, which comprises the steps of: (i) constructing a gene library of the protein of interest; (ii) preparing a vector by linking the gene library to a gene encoding spore coat protein; (iii) transforming a spore-forming host cell with the vector; (iv) forming a spore in the transformed host cell and displaying the protein of interest on a surface of the spore; (v) treating the spore displaying on its surface the protein of interest with one or more selected from the group consisting of organic solvent, heat, acid, base, oxidant, dryness, recovering the protease; (vi) and surfactant displaying on its surface the protein of interest; and (vii) screening the spore displaying a variant of the protein of interest having a resistance to the treatment.

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In further aspect of this invention, there is provided a method for bioconversion, which comprises the steps of: (i) preparing a vector for spore surface display comprising a gene construct containing a gene encoding spore coat protein and a gene encoding a protein of interest conducting a bioconversion reaction, wherein, when expressed, the gene construct expresses a fusion protein between the spore coat protein and the protein of interest; (ii) transforming a host cell with the vector for spore surface display; (iii) displaying the protein of interest on a surface of a spore of the host cell; (iv) recovering the spore displaying on its surface the protein of interest; and (v) performing the bioconversion reaction using the spore displaying on its surface the protein of interest.

In still further aspect of this invention, there is provided a method for preparing protein microarray, which comprises the steps of: (i) preparing a vector for spore surface display comprising a gene construct containing a gene—encoding spore coat protein and a gene encoding antibody or antigen having binding affinity to a protein to be analyzed, wherein, when expressed, the gene construct expresses a fusion protein between the spore coat protein and the antibody or antigen; (ii) transforming a host cell with the vector for spore surface display; (iii) displaying

the antibody or antigen on a surface of a spore of the host cell; (iv) recovering the spore displaying on its surface the antibody or antigen; and (v) immobilizing onto a solid surface the spore displaying on its surface the antibody or antigen.

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In another aspect of this invention, there is provided a method producing an antibody to antigen in vertebrates, which comprises the steps of: (i) preparing a vector for spore surface display comprising a gene construct containing a gene encoding spore coat protein and a gene encoding the wherein, when expressed, the gene construct antigen, expresses a fusion protein between the spore coat protein and the antigen; (ii) transforming a host cell with the vector for spore surface display; (iii) displaying the antigen on a surface of a spore of the host cell; (iv) recovering the spore displaying on its surface the antigen; composition vertebrates administering to and (\mathbf{v}) a containing an immunologically effective amount of the spore displaying on its surface the antigen.

In still another aspect of this invention, there is provided a method for preparing a whole cell absorber, which comprises the steps of: (i) preparing a vector for spore surface display comprising a gene construct containing a

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gene encoding spore coat protein and a gene encoding a protein having a binding affinity to a certain substance, wherein, when expressed, the gene construct expresses a fusion protein between the spore coat protein and the protein; (ii) transforming a host cell with the vector for spore surface display; (iii) displaying the protein on a surface of a spore of the host cell; (iv) recovering the spore displaying on its surface the protein; and (v) immobilizing onto a carrier the spore displaying on its surface the protein.

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According to preferred embodiments of this invention, the gene encoding spore coat protein is derived from a spore-forming Gram negative bacterium including Myxococcus; a spore-forming Gram positive bacterium including Bacillus; a spore-forming Actionmycete; a spore-forming yeast including Saccharomyces cerevisiae, Candida and Hansenulla or a spore-forming fungus, but not limited to. More preferably, the gene encoding spore coat protein is derived from a spore-forming Gram positive bacterium, most preferably, Bacillus including Bacillus subtilis and Bacillus polymyxa, etc.

The gene of spore coat protein useful in this invention includes cotA, cotB, cotC, cotD (W. Donovan et al., J. Mol. Biol., 196:1-10(1987)), cotE (L. Zheng et al., Genes & Develop., 2:1047-1054(1988)), cotF (S. Cutting et al., J.

Bacteriol., 173:2915-2919(1991)), cotG, cotH, cotJA, cotJC, cotK, cotL, cotM, cotS, cotT (A. Aronson et al., Mol. Microbiol., 3:437-444(1989)), cotV, cotW, cotX, cotY, cotZ (J. Zhang et al., J. Bacteriol., 175:3757-3766(1993)), spoIVA, spoVID and sodA, but not limited to.

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In addition, the gene encoding spore coat protein useful in this invention is a modified form or a recombinant form of one selected from the group consisting of cotA, cotB, cotC, cotD, cotE, cotF, cotG, cotH, cotJA, cotJC, cotK, cotL, cotM, cotS, cotT, cotV, cotW, cotX, cotY, cotZ, spoIVA, spoVID and sodA, in which the modified form recombinant form has a more compatibility for spore surface display relative to wild type genes. The modified form of the gene is obtained by DNA shuffling method (Stemmer, Nature, 370: 389-391(1994)), StEP method (Zhao, H., et al., Nat. Biotechnol., 16: 258-261 (1998)), RPR method (Shao, Z., et al., Nucleic acids Res., 26: 681-683 (1998)), molecular breeding method (Ness, J. E., et al., Nat. Biotechnol., 17: 893-896 (1999)), ITCHY method (Lutz S. and Benkovic S., Current Opinion in Biotechnology, 11: 319-324 (2000)), error prone PCR (Cadwell, R. C. and Joyce, G. F., PCR Methods Appl., 2: 28-33 (1992)), point mutagenesis (Sambrook et al., Molecular Cloning: A Laboratory Manual, Cold Spring Harbor, N. Y., 1989), nucleotide mutagenesis (Smith M. Annu. Rev. (1985)), combinatorial cassette 423-462 Genet. 19:

mutagenesis (Wells et al., Gene 34: 315-323 (1985)) and other suitable random mutagenesis.

Further to this, the gene encoding spore coat protein is selected from the group consisting of cotA, cotB, cotC, cotD, cotE, cotF, cotG, cotH, cotJA, cotJC, cotK, cotL, cotM, cotS, cotT, cotV, cotW, cotX, cotY, cotZ, spoIVA, spoVID and sodA, in which the gene has a substituted promoter for its promoter to enhance spore surface display relative to wild type genes. The promoter for enhancing surface display, for example, includes the promoters of cotE or cotG genes, which show higher expression level.

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In preferred embodiments of this invention, the gene encoding spore coat protein is selected from the group consisting of cotA, cotE, cotF, cotG, cotH, cotJA, cotJC, cotK, cotL, cotM, cotS, cotT, cotV, cotW, cotX, cotY, cotZ, spoIVA, spoVID and soda, more preferably, cotE or cotG and most preferably, cotG.

According to the present methods, as linking a gene of coat protein and a gene of the protein of interest, the overall sequence, fragments, two or more repeated sequences of the gene of coat protein are useful. In two or more repeated sequences, the repeated sequences may be the same or different each other. The overall sequence, two or more repeated sequences of the gene of the protein of interest are also useful in the fusion sequence. In two or more

repeated sequences, the repeated sequences may be the same or different each other. Other combinations also may be useful in the fusion sequence.

It is understood by one skilled in the art that the gene construct may exist as plasmid in host cell independently or as integrated form into chromosome of host cell. Additionally, in the gene construct, it is recognized by one skilled in the art that the gene of coat protein may be followed or preceded by the gene of the protein of interest. Integrated form into the counterpart gene may be useful.

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It is recognized by one skilled in the art that the expression of the fusion protein between coat protein and protein of interest can be induced by virtue of promoters of coat protein gene and protein of interest or other suitable promoters inducible in host cell

The present methods is applicable to any protein, for example, including enzyme, enzyme inhibitor, hormone, hormone analogue, hormone receptor, signal transduction protein, antibody, monoclonal antibody, antigen, attachment protein, structural protein, regulatory protein, toxin protein, cytokine, transcription regulatory protein, blood clotting protein, plant defense-inducing protein and fragments thereof. The applicable proteins include multimer as well as monomer. The surface display of multimeric proteins has been rarely reported, for instance, the surface

display of alkaline phosphatase in E. coli, has resulted the display toward inner part of cell outer membrane (Stathopoulus et al., 1996). β -galactosidase used as reporter enzyme in Examples of the present invention must form tetramer to exhibit its activity and has not been published to be successful in surface display. β galactosidase generally cannot pass across cell membrane and comprises an amino acid sequence detrimental to cell membrane, as a result, the fusion protein between surface display motif and β -galactosidase has been recognized not to be displayed on cell surface. Therefore, the surface display of β -galactosidase described in Examples proves to be surprising.

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The term used herein "protein" refers to molecule consisting of peptide bond, for example including oligopeptide and polypeptide.

The host cell suitable in this invention, includes sporeforming Gram negative bacterium including Myxococcus, a
spore-forming Gram positive bacterium including Bacillus, a
spore-forming Actionmycete, a spore-forming yeast and a
spore-forming fungus, but not limited to. Preferably, the
host cell is a spore-forming Gram positive bacterium, more
preferably, Bacillus. In particular, Bacillus subtilis is
advantageous in the senses that genetic knowledge and
experimental methods on its spore forming as well as

culturing method are well known.

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According to the present methods, the spore may be non-reproductive. In the method reproductive or improving a protein, the recovered coats are subject to reproduction but the methods using a spore as delivery means interest obviate the of necessity protein of reproduction of spore. It is considerable that the organisms genetically engineered is likely to be regulated under laws and rules; hence non-reproductive spore is preferable. For example, Bacillus subtilis lack of cwlD gene is preferably used due to being non-reproductive.

According preferred embodiments of this invention, the recovery of spore is performed in such a manner that the display of the protein of interest on the spore surface is maximized by controlling culture time, after which culturing is terminated and the spore is then recovered. Suitable culture time is varied depending upon the type of cell used, for example, in case of using *Bacillus subtilis* as host, the culture time of 16-25 hours is preferred.

In the present methods, the recovery of spore may be carried out according to the conventional methods known to one skilled in the art, more preferably, renografin gradients methods (C. R. Harwood, et al., "Molecular Biological Methods for Bacillus." John Wiley & Sons, New York, p.416(1990)).

As demonstrated in Examples, the stability of spore displaying the foreign protein of interest on its surface is very high in the present invention, indicating maintenance of the integrity of spore surface structure formed by cooperation of coat proteins while the foreign protein is displayed.

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The protein of interest displayed on spore surface according to the present methods can be demonstrated with a wide variety of methods as follows: 1) A primary antibody is bound to the protein of interest displayed on spore surface and then reacted with a secondary antibody labeled with chemical to stain the fluorescent spore, followed by observation with fluorescence microscope or analysis with flow cytometry. 2) The protein of interest displayed on spore surface is treated with protease, followed by measurement of the activity of the protein or detecting lower signal with fluorescence microscope or flow cytometry. 3) In case that the protein of interest uses a substrate with higher molecular weight, the direct measurement of the activity of the protein can provide the level of display since the substrate cannot pass across outer coat of spore.

In the method for improving protein, the construction of gene library for the protein of interest is performed by a mutagenesis of the gene encoding the protein of interest of

wild type, in which the mutagenesis includes DNA shuffling method (Stemmer, Nature, 370: 389-391(1994)), StEP method (Zhao, H., et al., Nat. Biotechnol., 16: 258-261 (1998)), RPR method (Shao, Z., et al., Nucleic acids Res., 26: 681-683 (1998)), molecular breeding method (Ness, J. E., et al., Nat. Biotechnol., 17: 893-896 (1999)), ITCHY method (Lutz S. and Benkovic S., Current Opinion in Biotechnology, 11: 319-324 (2000)), error prone PCR (Cadwell, R. C. and Joyce, G. F., PCR Methods Appl., 2: 28-33 (1992)), point mutagenesis (Sambrook et al., Molecular Cloning: A Laboratory Manual, Cold Spring Harbor, N. Y., 1989), nucleotide mutagenesis (Smith Genet. 19: 423-462 (1985)), Μ. Annu. Rev. combinatorial cassette mutagenesis (Wells et al., Gene 34: 315-323 (1985)) and other suitable random mutagenesis.

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In the method for improving protein, the screening is performed in a rapid manner by means of measuring an activity of the protein or flow cytometry (Georgiou, 2000). In case of using an activity of the protein, the screening is carried out by measuring growth of host expressing the protein or colorimetrical reaction catalyzed by the protein. In the method for improving protein using a resistance property of spore, the screening is carried out in a rapid manner by virtue of measuring an activity of the protein or using the structural stability of the protein.

The methods for improving protein provide in a high-

throughput manner, from wild type, 1) enzymes catalyzing non-biological reaction (e.g., Diels-Alder condensation), 2) enzymes with non-natural steroselectivity or regioselectivity, 3) enzymes with activity in organic solvent or organic solvent-aqueous solution two-phase system, and 4) enzymes with activity in extreme conditions such as high temperature or pressure.

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In addition, to select a variant of antibody with enhanced binding affinity, it is general that pH is abruptly changed or the concentration of base is adjusted to elute the variant. In a method using phage or bacteria as carrier, such elution conditions are likely to decrease the viability of phage or bacteria in medium. However, the methods for improving protein using system of spore surface display overcome the drawback.

In the meantime, the bioconversion process using surface-displayed enzymes requires a physiochemical stability of surface displaying host in extreme conditions because the process is usually executed in high temperature and/or organic solvent. In particular, a chemical synthesis valuable in current industry is mainly carried out in organic solvent and the synthesis of chiral compound or the resolution of racemic mixture is also performed in highly severe physiochemical conditions. Therefore, the surface-displayed enzyme as well as the organisms displaying enzyme

is compelled to have stability in such extreme conditions. In this connection, it is demonstrated that the methods for bioconversion using system for spore surface display is largely advantageous.

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The chemical processes using surface-displayed enzymes have been proposed (Georgiou et al., 1993). However, the proposed processes have generally required immobilization of surface with cross-linking agent cell since the displaying enzyme is very unstable during process (Freeman et al., 1996). The present bioconversion process is free from the disadvantage mentioned above. Because the surfacedisplayed enzyme as well as the host displaying enzyme is largely stable, the present method avoids the immobilization. Examples described hereinafter, In the bioconversion reaction with β -galactosidase is exemplified and thus it is understood by one skilled in the are that the present method can be also applied to any type of enzyme such as lipase, protease, cellulase, glycosyltransferase, oxidoreductase and aldolase. In addition, the present method is useful in single step or multi-step reaction and in aqueous or nonaqueous solution. The present bioconversion method employs spore as free or immobilized form and can be performed with other microbes or enzymes.

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Similar to DNA microarray, a protein microarray provides means for analyzing expression or expression level of target

protein in certain cell. In order to fabricate protein array, the suitable proteins to be arrayed must be obtained and then immobilized on solid surface. During analysis using protein array, washing step is necessarily performed to remove unbound proteins and various treatments such as high temperature, higher salt concentration and pH adjustment are executed; therefore, it is pivotal to guarantee proteinaceous substance with higher stability in such detrimental environment.

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In addition, the conventional process for preparing protein array needs tedious and repetitive works such as cloning genes of several thousands to tens of thousands of proteins and immobilizing of the proteins expressed. Therefore, there remains a need to improve simplicity and rapidity of the works.

According to the method for preparing protein microarray of this invention, it is ensured that the works described-above cane be performed with much greater readiness. In the present method, a gene construct containing a gene encoding spore coat protein and a gene encoding the desired protein is introduced into host cell and the spore displaying on its surface the desired protein is isolated, followed by immobilization of the isolated spore onto a solid surface. In the method for preparing protein array, the conventional steps may be used (see WO 0061806, WO 0054046, US 5807754,

EP 0818467, WO 9742507, US 5114674 and WO 9635953). The protein microarray manufactured by the present invention has a variety of applicable fields including diagnosis, analysis of gene expression, analysis of interaction between proteins, analysis of interaction between protein and ligand, study on metabolism, screening novel or improved enzymes, combinatorial biochemical synthesis and biosensor.

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solid substrate suitable in the present method includes, but not limited to, glasses (e.g., functionalized glasses), Si, Ge, GaAs, GaP, SiO, SiN4, modified silicone nitrocellulose, polyvinylidene fluoride, polystylene, polytetrafluoroethylene, polycarbonate, nylon, fiber and combinations thereof. The spore optionally may be attached the array substrate through linker molecules. It is preferred that the regions of the array surface not being spotted are blocked. The amount of spores applied to each spot (or address) depends on the type of array. Interaction between the protein displayed on spore attached to solid substrate and the sample applied can be detected based on their inherent characteristics (e.g., immunogenicity) or can being labeled with detectable rendered by an be independently detectable tag (e.g., fluorescent, luminescent or radioactive molecules, and epitopes). The data generated with protein array of this invention can be analyzed using known computerized systems such as "reader" and "scanner".

According to the method producing an antibody of this invention, a composition containing an immunologically effective amount of the spore, preferably, further comprises adjuvant such as incomplete and complete Freund's adjuvants. In the present method, the mode of administration is, preferably, injection and more preferably, intravenous, intraperitoneal, subcutaneous and intramuscular injections. within Boosting suitable period after the first administration is preferable to yield a sufficient amount of antibody.

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Meanwhile, in the process for preparing absorption chromatography, antibody or polypeptide is produced, purified and immobilized on a carrier. Generally, it is very difficult to prepare the bioabsorbers. The disadvantage may be overcome using whole cell displaying protein as described in Georgiou et al., 1997. Therefore, the system for spore surface display of this invention provides a whole cell absorber to solve the problems of the known absorbers.

In further aspect of this invention, there is provided a microbial transformant for spore surface display of a protein of interest, characterized in that the transformant is produced by transformation with a vector for spore surface display containing (i) a gene encoding a protein of interest and (ii) a gene encoding spore coat protein is

selected from the group consisting of cotA, cotB, cotC, cotD, cotE, cotF, cotG, cotH, cotJA, cotJC, cotK, cotL, cotM, cotS, cotT, cotV, cotW, cotX, cotY, cotZ, spoIVA, spoVID and sodA, in which when expressed, a fusion protein between the spore coat protein and the protein of interest is expressed.

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According to preferred embodiment, the transformant is derived from a variant mutated to enhance spore surface display. For example, the mutation to enhance spore surface display eliminates a production of extracellular secretory protease in the transformant, so that the protein of interest displayed on spore surface is stably maintained. In addition, the mutation to enhance spore surface display eliminates a production of intracellular protease in the transformant. It is also preferred that a gene or genes involved in spore forming is subject to mutation in order to the rate of spore forming (Perego, M., et al., Mol. Microbiol. 19: 1151-1157 (1996)).

In still further aspect of this invention, there is provide a spore for spore surface display of a protein of interest, characterized in that the spore displays the protein of interest on its surface.

According to the present invention, the spore may be reproductive or non-reproductive one which is selected based on its application field. Preferably, the non-reproductive

spore can be obtained by virtue of one or more methods selected from the group consisting of genetic method (Popham D. L., et al., J. Bacteriol., 181: 6205-6209 (1999)), chemical method (Setlow T. R., et al., J. Appl. Microbiol., 89: 330-338 (2000)) and physical method (Munakata N, et al., Photochem. Photobiol., 54: 761-768 (1991)). The genetic method to make the spore non-reproductive is accomplished by, for example, deleting a gene of host cell involved in reproduction of spore.

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In the present invention, it is preferred that the spore is derived from a variant mutated to increase its agglutination property because in bioconversion performed in industrial scale, the separation between the resulting product and spores is rendered easier. The increase of the agglutination property in the spore is accomplished by one or more methods selected from the group consisting of genetic method, chemical method and physical method. As example of the physical method, the heat treatment can be proposed (Wiencek K. M., et al., Appl. Environ. Microbiol.,

In another aspect of this invention, there is provided a vector for spore surface display, characterized in that the vector comprises a replication origin, an antibiotic-resistance gene, a restriction site, a gene encoding a spore

coat protein, a gene encoding a protein of interest and a promoter operatively linked to the gene encoding a spore coat protein, in which when expressed, a fusion protein between the spore coat protein and the protein of interest is expressed.

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According to preferred embodiment, the gene encoding a spore coat protein is selected from the group consisting of cotA, cotB, cotC, cotD, cotE, cotF, cotG, cotH, cotJA, cotJC, cotK, cotL, cotM, cotS, cotT, cotV, cotW, cotX, cotY, cotZ, spoIVA, spoVID and sodA, more preferably, cotE or cotG, and most preferably, cotG.

In the vector of this invention, the replication origin can include various origins known to one skilled in the art, for example, when the vector is introduced into a sporeforming yeast, 2µ, ARS, ARS1 or ARS2 can be used as replication origin. In case of using Bacillus as host, ori ColEl origin, Rep1060, etc. can be used. The antibiotic-resistance gene used as selective marker, when prokaryote such as Bacillus is used as host, is a resistance gene to antibiotics acting to prokaryotes, for example, carbenicillin, kanamycin, ampicillin, including chloramphenicol, streptomycin, geneticin, neomycin tetracycline. The promoter used in the present vector includes a promoter of the gene of spore coat protein and a known promoter operable in host cell.

BRIEF DESCRIPTION OF THE DRAWINGS

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Fig. 1 is a microscopic photograph showing spores of Bacillus subtilis purified according to method described in U.S. Pat. No. 5,766,914;

- Fig. 2 is a microscopic photograph showing spores of Bacillus subtilis purified according to renografin gradients method;
- Fig. 3 is a genetic map of the recombinant vector pCotE
 lacZ of the present invention;
 - Fig. 4 is a genetic map of the recombinant vector pCotG-lacZ of the present invention;
 - Fig. 5 represents screening results demonstrating the preferred surface display motif in the present invention;
- Fig. 6 is a graph showing the affect of protease to β galactosidase displayed on spore surface;
 - Fig. 7 is a graph showing the activity of β -galactosidase displayed on spore surface in accordance with culture time;
- Fig. 8 is a graph representing the heat stability of Bacillus subtilis DB104 strain displaying on its surface the protein;
 - Fig. 9 is a genetic map of recombinant vector pCSK-cotG-CMCase of this invention;
- Fig. 10 is a graph showing analysis of spore surface-

displayed carboxymethylcellulase using flow cytometry;

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Fig. 11 is a graph showing analysis of spore surface-displayed levansucrase using flow cytometry;

Fig. 12 is a graph showing the activity of spore surface-displayed levansucrase;

Fig. 13 is a graph representing analysis of spore surface-displayed monoclonal antibody using flow cytometry;

Fig. 14 is a graph demonstrating selectivity to spore displaying single chain Fv;

Fig. 15 is a graph representing analysis with flow cytometry of monoclonal antibody library to have binding affinity to Pre-S region of hepatitis B virus;

Fig. 16 is a graph showing analysis of spore surface-displayed GFP using flow cytometry; and

Figs. 17a to 17d are graphs representing isolation with flow cytometry of spores displaying improved GFP.

The following specific examples are intended to be illustrative of the invention and should not be construed as limiting the scope of the invention as defined by appended claims.

EXAMPLES

Example I: Isolation of the Gene Encoding Coat Proteins

I-1: Construction of the Vector for Spore Surface Display

To isolate the most appropriate coat protein for spore surface display among coat proteins consisting of spore, the recombinant vector having the gene encoding a fusion protein between coat protein and β -galactosidase was constructed as follow:

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To begin with, the DNA was extracted from the Bacillus subtilis 168 strain provided from Dr. F. Kunst (Kunst F., et al., Nature, 390: 249-256(1997)) by Kalman's method (Kalman S., et al., Appl. Environ. Microbiol. 59, 1131-1137(1993)), and the purified DNA was served as template for PCR to spoIVA primers (SEQ ID NOs: 1 and 2), cotB primers (SEQ ID NOs: 3 and 4), cotC primers (SEQ ID NOs: 5 and 6), cotD primers (SEQ ID NOs: 7 and 8), cotE primers (SEQ ID NOs: 9 and 10), cotG primers (SEQ ID NOs: 11 and 12), cotH primers (SEQ ID NOs: 13 and 14), cotM primers (SEQ ID NOs: 15 and 16), cotV primers (SEQ ID NOs: 17 and 18), cotX primers (SEQ ID NOs: 19 and 20) and coty primers (SEQ ID NOs: 21 and 22). Tag polymerase purchased from Boehringer Mannheim was used for total 35 cycles of PCR under condition of denaturation for 30 sec at 94°C, annealing for 30 sec at 55°C and extension for 1 min at 72° C.

After then, each amplified PCR products were digested with BamHI and SalI and cloned between BamHI and SalI sites of plasmid pDG1728 which is a gratuitous gift by Dr. P. Stragier (Geurout-Fleury, A.M., et al., Gene, 180: 57-

61(1996)), thus the constructed vectors express the fusion protein of coat protein and β -galactosidase. Fig. 3a shows the genetic map of pCotE-lacZ expressing fusion protein of CotE protein and β -galactosidase and Fig. 3b shows the genetic map of pCotG-lacZ expressing fusion protein of CotG protein and β -galactosidase.

SEQ ID NO:23 shows the sequence of cotE-lacZ fused genes and SEQ ID NO:24 shows the amino acid sequence of CotE-LacZ fusion protein. In SEQ ID NO:23, promoter for cotE is 1-329, CotE structural gene is 330-872, restriction site is 873-878 and LacZ structural gene is 879-3902.

SEQ ID NO:25 shows the sequence of cotG-lacZ fused genes and SEQ ID NO:26 shows the amino acid sequence of CotG-LacZ fusion protein. In SEQ ID NO:25, promoter of cotG is 1-460, CotE structural gene is 461-1045, restriction site is 1046-1051 and LacZ structural gene is 1052-4075.

I-2: Pure Isolation of Spores

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Constructed recombinant expression vectors were transformed into Bacillus subtilis DB104 (Kawamura F. and Doi R.H., J. Bacteriol. 160: 442-444(1984)) using natural transformation (C.R. Harwood, et al., Molecular Biological Methods for Bacillus, John Wiley & Sons, New York, p.416(1990)).

Other methods such as conjugation or trnasduction can be

applied for introduction of the recombinant vectors into Bacillus strain.

Subsequently, each *Bacillus* strain comprising the fused gene between coat protein and β -galactosidase was cultured for 24 hr at a shaking incubator (37°C, 250 rpm) in GYS medium ((NH₄)₂SO₄ 2 g/l, Yeast extract 2 g/l, K₂HPO₄ 0.5 g/l, glucose 1 g/l, MgSO₄D5H₂O 0.07 g/l), and the only pure spores were isolated using renografin gradients method (C. R. Harwood, et al., "Molecular Biological Methods for Bacillus." John Wiley & Sons, New York, p.416(1990)).

I-3: Display of Proteins on Spore Surface

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The spores isolated in the above-described Example and the cell pellet of Bacillus subtilis DB104 were subjected to evaluation of the activity of β-galactosidase using Miller's method (Miller, "Experiments in Molecular Genetics", Cold Spring Harbor, NY: Cold Spring Harbor Laboratory, p.352-355(1972)) and the results are shown in Fig. 5. In Fig. 5, the gray bar indicates cell pellet, the black bar indicates the activity of β-galactosidase in purely isolated spores and '1' relates to result of control Bacillus subtilis DB104; '2' to result of SpoIVA-LacZ; '3' to result of CotB-LacZ; '4' to result of CotC-LacZ; '5' to result of CotD-LacZ; '6' to result of CotE-LacZ; '7' to result of CotG-LacZ; '8' to result of CotM-LacZ; '9' to result of CotM-LacZ; '8' to result of CotM-

LacZ; '10' to result of CotV-LacZ; '11' to result of CotX-LacZ; and '12' to result of CotY-LacZ fusion protein, respectively.

As shown in Fig. 5, it is known that Deits TL (U.S. Pat. No. 5,766,914) fails to induce the sufficient surface display of cotC and cotD since the expression levels of cotC and cotD are as low as the control. However, the expression level of cotE and cotG are comparatively high and especially, expression level of cotG is remarkably high comparing to other coat proteins. In addition, in the isolated spores, the surface display using cotG shows the highest enzyme activity, which demonstrates that CotG-LacZ fusion proteins are the highest level of display on spore surface.

Considering the expression level and the amount of fusion proteins displayed on spore surface, it is known that the cotG is the most preferable surface display motif. It is known to one skilled in the art that these results exclude other coat proteins other than cotG from applying to spore surface display.

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I-4: Effect of Proteases on the Surface-Displayed Enzymes

To confirm whether the surface-displayed β -galactosidase is degraded or not, the purely isolated spore displaying CotG-LacZ was resuspended into 100 μ l of PBS solution, and

each 10 mg/ml of protease K, protease type XIV or trypsin was treated. Thereafter, the activity of β -galactosidase was measured as described above and the results are shown in Fig. 6. As shown in Fig. 6, the activity of spore surfacedisplayed β -galactosidase is decreased with some variations in each result. These results give the evidence for the localization of β -galactosidase on spore surface.

DB104 strain lacking neutral and alkaline protease and (Ye, R., et al., Biotechnology strain WB700 Bioengineering, 62:87-96(1999)) lacking 7 proteases among proteases secreted from Bacillus subtilis were transformed pCotG-lacZ expression vector using natural with the transformation method as described in example I-1, and the activity of β -galactosidase in cell pellet and spores was measured as described in example I-3 (Fig. 7). As shown in Fig. 7, while the enzyme activity is abruptly decreased in DB104 strain as time goes, WB700 strain shows slight decrease in enzyme activity. These results indicate that the displayed \beta-galactosidases on spore surface are degraded in DB104 strain by the proteases secreted extracellularly; however, the displayed β -galactosidases in WB700 are stably maintained because of lack of the proteases secreted extracellularly. Therefore, the results also support the localization of β -galactosidase on spore surface.

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Example II: Spore Production Depending on Culture Time

As shown in Fig. 7, it is required to stop incubation on a specific time point and isolate spores. In DB104, the enzyme activity of spores after 38 hr of incubation is significantly low comparing to that after 24 hr of incubation. Thus, it is demonstrated that the adjustment of incubation time makes it possible to yield spores displaying enzyme on its surface with the greatest enzyme activity.

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Example III: Characterization of Spores Dispalying β galactosidase

follow in measured as spores resistance was Heat displaying β -galactosidase: 100 μ l of spores isolated by renografin gradients in Example I-2 were heated for 15 min and then spread on LB plates to evaluate viability of spores (Fig. 8). As shown in Fig. 8, spores displaying CotG-LacZ show similar heat resistance to spores without surface protein. In a result, the display of the foreign protein fused to coat protein on spore surface does not affect on inherent characteristics such as heat resistance. its Moreover, these results provide the promising usage of spore displaying on its surface enzyme in chemical reactions at high temperature. In addition, from these results, it is suggested that the spores transformed according to the present invention remain their inherent resistances to

lysozyme, a bacterial cell wall-degrading enzyme and solvent.

Example IV: Displaying Various Enzymes on Spore Surface

IV-1: Construction of Recombinant Vectors

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spores displaying various enzymes, To use it is prerequisite to confirm whether various enzymes in addition β-galactosidase can be surface-displayed. Firstly, to plasmid pHPS9 (Haima, et al., Gene, 86:63-69(1990)) was digested by EcoRI and HindIII and manipulated into blunt ends using Klenow enzyme. Then, DNA fragment containing multiple cloning sites, which was obtained from plasmid p123T (EMBL Z46733) with BssHII, was ligated to the bluntended pHPS9 plasmid to use as virgin vector named pCSK1 in following experiments. The pCSK-cotG plasmid the prepared by restricting pCSK1 plasmid with BamHI and PstI and ligating PCR-amplified cotG gene. In the course of PCR for cotG gene amplification, a linker between cotG gene and target gene was incorporated using cotG-linker 5 primer (SEQ ID NO:27) and 3 primer (SEQ ID NO:12) with template of DNA in Bacillus subtilis.

In other experiments, genes encoding carboxymethyl cellulase, levansucrase and lipase was prepared as follows: Carboxymethyl cellulase cloned in pBSI plasmid (S. H. Park et al., Agric. Biol. Chem., 55: 441-448(1991)) was directly

employed. The pBS1 plasmid contains the gene encoding carboxymethylcelluase cloned from Bacillus subtilis BSE616 strain. In the present Example, PCR was performed with the pBS1 as template using primer represented by SEQ ID NOs:28 and 29. In the case of PCR for levansucrase, pSSTS110 plasmid (Jung, H.-C., et al., Nat. Biotech., 16; 576-580(1998)) was used as template and primers represented by SEQ ID NOs:30 and 31 were used. In PCR for lipase, pTOTAL (Ahn, J.-H., et al., J. Bacteriol., 181: 1847-1852(1999)) was added as template and primers of SEQ ID NOs: 32 and 33 were used. All PCRs were performed in the same condition as described in Example I-1.

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Recombinant vectors containing gene coding for fusion between CotG and the carboxymethylcelluase, levansucrase or lipase were prepared by cloning into pCSK-cotG using PstI and BamHI restriction enzymes both in vector and in the PCRamplified inserts. As an example of the above construction, shows pCSK-cotG-CMCase which is the recombinant vector fusion protein encoding between CotG and carboxymethylcellulase. Transformed Bacillus subtilis DB104 with pCSK-cotG-CMCase was named Bacillus subtilis GFSD18 and deposited at the Korean Collection for Type Cultures (KCTC, KR) with accession No. KCTC 0887BP (November 16, 2000).

SEQ ID NO:34 shows nucleotide sequence of fused cotG-CMCase genes and SEQ ID NO:35 shows amino acid sequence of

CotG-CMCase encoded by SEQ ID NO:34. In SEQ ID NO:34, promoter for cotG is 1-460, structural gene for CotG is 461-1045, linker is 1046-1084, and structural gene for CMCase is 1085-2491.

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IV-2: Expression of Recombinant Vectors and Verification

The above-prepared recombinant vectors were employed for transformation of Bacillus subtilis DB104 with the same procedures as described in Example I-2. Subsequently, each transformed Bacillus strains was cultured for 24 hr at a shaking incubator (37°C, 250 rpm) in GYS midium, the only pure spores were isolated using renografin gradients method, and enzyme activity of carboxymethylcellulase (Kim, et al., Appl. Environ. Microbial., 66:788-793(2000)), levansucrase (Jung, et al., Nat. Biotech., 16:576-580(1998)) or lipase was evaluated. The activity of lipase was evaluated as follow: The spores suspended in 10% PBS was mixed with 10% olive oil, reacted for 48 hr, treated with 0.2 ml cupric acid on supernatant solution and the observance of OD was performed at 715 nm.

In the case of carboxymethylcellulase, the activity of enzyme displayed on spore was 175 mU comparing to 0 mU in control. In other verifying method, carboxymethylcellulase-specific andtibody (Kim, et al., Appl. Environ. Microbiol., 66:788-793(2000)) was probed for flow cytometry (FACSort,

Becton Dickinson, USA) and the carboxymethylcellulases were detected on the surface of spores transformed by pCSK-cotG-CMCase (Fig. 10).

The activity of levansucrase was also high in spores transformed by recombinant vector (Fig. 12) and the levansucrases were detected on the surface of transformed spores as verified with flow cytometery using levan sucrase-specific antibody (Jung, et al., Nat. Biotech., 16:576-580(1998)) in the same procedures as above-described in carboxymethylcellulase (Fig. 11).

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The activity of lipase was measured as $A_{715} = 0.14$ in spores transformed with recombinant vector.

On the basis of these results, it is demonstrated that various enzymes as well as β -galactosidase can be displayed on the surface of spore according to the present invention.

Based on the results in these examples and example I, it is known to one skilled in the art that the gene construct containing gene encoding fusion protein between coat protein and protein of interest may exist as plasmid in host cell independently or as integrated form into chromosome of host cell and both forms may lead to successful spore surface display. It is also recognizable that the gene of coat protein may be followed or preceded by the gene of the protein of interest. In addition, it is recognized that in

the gene construct, the overall sequence, fragments, two or more repeated sequences of the gene of coat protein are useful. In two or more repeated sequences, the repeated sequences may be the same or different each other. The overall sequence, two or more repeated sequences of the gene of the protein of interest are also useful in the fusion sequence. In two or more repeated sequences, the repeated sequences may be the same or different each other.

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It is recognized by one skilled in the art that the expression of the fusion protein between coat protein and protein of interest can be induced by virtue of promoters of coat protein gene and other suitable promoters operable in host cell. Any vector carrying the present gene construct may be used in this invention, which is recognized by one skilled in the art referring to these results.

It is known that both monomric and multimeric enzyme can be applied for the present invention since the β -galactosidase used in example I is tetramer (U. Karlsson et al., J. Ultrastruct. Res., 10:457-469(1964)) and the enzymes described in this Example are monomers.

Example V: Display of Antibody on Spore Surface and Screening for Directed Evolution

On the purpose of application of other proteins in addition to enzymes, the experiment to display antibody on

spore surface was performed as follows:

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V-1: Construction of Recombinant Vector for Surface Display of Single Chain Fv

Gene encoding single chain Fv against Pre-S2 domain (SEQ ID NO:36) of hepatitis B virus (HBV) was linked to cotG gene encoding surface protein of Bacillus subtilis spore. Single chain Fv gene was amplified by PCR with pAScFv101 (WO 9737025) as template and with primers described in SEQ ID NOs:37 and 38. Tag polymerase purchased from Bioneer (Korea) was used for total 30 cycles of PCR under condition of denaturation for 30 sec at 94 °C, annealing for 30 sec at 55 $^{\circ}$ C and extension for 1 min at 72 $^{\circ}$ C. And then, each PCR product was restricted by ApaI and NheI, cloned into pCSK-CotG between the same restriction sites (pCSK-CotG-scFv) and transformed into JM109 using transformation method by Inoue, et al. (Inoue, H., et al., Gene, 96:23-28(1990)). The amplified vectors for displaying on spore surface were isolated by alkaline extraction method (Sambrook et al., Molecular Cloning: A laboratory Manual, Cold Spring Harbor, N.Y., 1989) and transformed into Bacillus subtilis DB104 by natural transformation as described in Example I-1.

V-2: <u>Verification of Single Chain Fv Display on Spore</u> Surface Using Flow Cytometry

Affinity of the displayed single chain Fv against the Pre-S2 of HBV was evaluated by FACSort as the following procedures.

Firstly, Pre-S2 peptide was labeled with fluorescein (PanVera, USA) using fluorescein succinidimyl ester coupling method.

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The transformed strains were inoculated into LB broth containing 5 μ g/ml chloroamphenicol, pre-cultured for 8-10 hr at 37°C, 1% of seed culture was inoculated into GYS broth for sporulation, cultured for 24 hr at 37°C and the cultured media was harvested. The pure spores were isolated using renografin gradients method, 100 μ l pure spores were blocked with PBS containing 3% skim milk to inhibit non-specific binding and reacted with 10 μ l of fluorescein labeled Pre-S2 peptide. Thereafter, the spores bound to fluorescein labeled Pre-S2 peptide were detected in the same procedures as described in example IV (Fig. 13). As shown in Fig. 13, it is demonstrated that the monoclonal antibody against Pre-S2 peptide is successfully displayed without reduction of the affinity to its antigen.

According to the above results, it is recognized that the present methods may be applicable to any protein, for example, enzyme, hormone, hormone analogue, enzyme inhibitor, signal transduction protein or its fragment, antibody or its fragment, antigen protein, attachment protein, structural

protein, regulatory protein, toxin protein, plant defenseinducing protein.

V-3: Selection of Spores Displaying Single Chain Fv using Flow Cytometry

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Whether the displayed single chain Fv has affinity to Pre-S2 of HBV was verified with FACSort as follows:

The transformed strains were inoculated into LB broth containing 5 μ g/ml chloroamphenicol, pre-cultured for 8-10 hr at 37°C, 1% of seed culture was inoculated into GYS broth for sporulation, cultured for 24 hr at 37°C and the cultured media was harvested. And then, 50 ml of harvested culture medium was centrifuged at 10,000 g for 10 min, supernatant was discarded, bacteria were resuspended in 500 μ l of 20% renografin (Sigma, USA). 100 μ l of resuspended cell was carefully flowed onto 500 μ l of 50% renogrant in microtube to form layer, the microtube was centrifuged at 10.000 g for 30 min and pure spores were isolated from pellet.

To discard remained renografin, spores were rinsed 3 times with DW and resuspended in PBS buffer. And then, spores displaying single chain Fv were mixed with wild type spores at a ratio of 1:103 and 1:105 and the spores with affinity to Pre-S2 of HBV were harvested using fluorescein-labeled Pre-S2 peptide and FACSort.

The selectivity was evaluated by colony-forming assay on

LB agar plates and LB agar plates containing 5 $\mu g/ml$ of chloroamphenical comparing to wild type. Spores displaying surface single chain Fv are resistant to chloroamphenical owing to chloroamphenical resistant gene contained in the recombinant vectors.

Fig. 14 shows the selectivity of spores displaying single chain Fv in each ratio (selectivity = ratio of spores displaying single chain Fv after flow cytometry/ ratio of spores displaying single chain Fv before flow cytometry). In the case that the ratio of spores displaying single chain Fv before flow cytometry is 10⁻⁵, the selectivity was over 4,000, which indicates that spores with enhanced affinity can be selected by flow cytometry among spores displaying various antibody libraries.

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V-4: Directed Evolution of Single Chain Fv Displayed on Spore Surface

To display single chain Fv library on spore surface, the gene encoding single chain Fv against Pre-S2 of HBV was amplified by error—prone PCR.—PCR was carried out using pAScFv101 plasmid described in the example V-1 as template and SEQ ID NOs:37 and 38 as primer. PCR mixture was prepared by mixing 0.3 µM of each primers, 5 ng of DNA template, PCR solution (10mM Tris(pH 8.3), 50 mM KCl, 7 mM MgCl₂, 0.01% (w/v) gelatin), 0.2 mM dGTP, 0.2 mM dATP, 1 mM dTTP, 1 mM

dCTP, 5 U Taq polymerase from Bioneer (Korea) and DW up to 100 μ L. Total 13 cycles of PCR was performed under condition of denaturation for 30 sec at 94°C, annealing for 30 sec at 50°C and extension for 1 min at 72°C.

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Subsequently, restricted PCR products with ApaI and NheI were cloned into pCSK-CotG, vector for displaying on spore surface, between the same restriction sites and library was prepared by transforming the cloned vectors into JM109 E. coli with the method of Inoue et al.

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The vectors for displaying on spore surface were isolated by alkaline extraction method and transformed into Bacillus subtilis DB104 by natural transformation. And then, single chain Fv library against Pre-S2 of HBV was displayed on spore surface as described in example V-2 (Fig. 15).

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As shown in Fig. 15, spores with increased fluorescence (i.e., increased affinity) were isolated. This result demonstrates the applicability of the present invention to prepare and select protein variants with improved characteristics.

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Example VI: Bioconversion using Spores Displaying Protein of Interest

Forte of transglycosylation by enzyme is the capability of formation of site-specific glycosidic linkage without protection/de-protection step. There have been studied for

formation of glycosidic linkage by 1) induction of reverse hydrolysis in non-aqueous system using glycosidase which is conventionally available glycosidic hydrolyzing enzyme and transglycosylation in which glycosidic linkage 2) substituted with receptor alcohol instead of hydrolysis of glycosidic linkage by water (G. Ljunger et al., Enzyme Microb. Technol., 16:1808-1814(1994); T. Usui et al., Carbohytdr. Res., 244:315-323(1993); and R. Lopez et al., J. Org. Chem., 59:737-745(1994)). The above conventional methods usually use organic solvent to increase synthetic yield and inhibit hydrolysis. However, because the organic solvent inactivates enzyme, it is difficult to accomplish the high yield. Thus, it is necessary to inhibit the inactivation of glycosidase in organic solvent for higher glycosylation yield.

The purpose of the Example is to exemplify the higher glycosylation yield with improved enzyme stability even in organic solvent by virtue of displaying glycosidase on the surface of hydrophobic *Bacillus* spores.

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VI-1: Stability of β -galactosidase Displayed on Surface of Spores in Organic Solvent

Each of β -galactosidase in free form (Sigma, USA) and the β -galactosidase displayed on surface of *Bacillus* spore was dispersed into 500 μ l of Tris-HCl buffer (pH 7.5), added the

same volume of the various solvents described in Table 1, mixed for 37° C for 1 hr and the remained enzymatic activity was measured by Miller method described in Example I-3 (Table 1).

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TABLE 1

TABLE 1		
	Residual activity (%)	
	Free form	Surface-Displayed
	β-galactosidase	β-galactosidase
Control	100	100
Hexane	84.3	100
Ether	48.2	77.2
Toluene	4.2	51.9
Ethylacetate	0.1	9.6
Acetonitril	0.0	0.8
Ethanol	0.0	0.0

As shown in Table 1, the displayed β -galactosidase shows higher stability than that of free form β -galactosidase in various organic solvents.

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VI-2: Transglycosylation Reaction in Water-Organic Solvent Two-phase System Using β -galactosidase Displayed on Spore Surface

To perform transglycosylation in two-phase system, β -galactosidase, which is one of conventional glycosidase, is

used as a model for glycosylation reaction (Scheme 1).

Scheme 1

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At first, 1 ml of 1 M lactose in 10 mM phosphate buffer (pH 5.1) was mixed with 10 ml of 10 mM 5-phenyl-1-pentanol in hexane for reaction solution. And then, β -galactosidase displayed on spore surface (240 U; 1 U = the amount of enzyme capable of hydrolysis of 1 μ mol ONPG (o-nitrophenyl- β -D-galactopyranoside) for 1 min at 37°C) and free form β -galactosidase (240 U) was added into the above reaction solution, respectively, and reacted for 48 hr at 30°C while stirring.

results, the yield of 5-phenylpenthyl-β-D-In galtopyranoside was 21% by β -galactosidase displayed on spore surface; however, in free form β-galactosidase, the hydrolysis of lactose only occurred with no transglycosylation. Such result is ascribed to the increased stability, in organic solvent, of β -galactosidase displayed on spore surface. Actually, after 72 hr reaction, about 5% enzyme activity was detected in the displayed β of

galactosidase while measured the complete inactivation in free form β -galactosidase. Another advantage of the displayed β -galactosidase owes to hydrophobicity of Bacillus spores. In other words, the distribution of displayed β -galactosidase at interface between water and organic solvent phase inhibits the hydrolysis comparing to free form β -galactosidase.

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Based on the results of this Example, it is understood that variants and modifications thereof falling within the spirit of the invention may become apparent to those skilled in the art, for example, any enzymes in addition to β -galactosidase such as lipase and protease can be employed for bioconversion of the present invention. In addition, the present bioconversion is useful in single step or multi-step reaction and in aqueous or non-aqueous solution. The present bioconversion method can employ spore as free or immobilized form and can be performed with other microbes or enzymes.

Example VII: Display of Antigen on Spore Surface

By displaying antigen on spore surface, antigen capable of inducing immune response in vivo can be applied as live vaccine. Bacillus subtilis has been considered as safe strain to human body since it has been employed in food fermentation for a long time (Sonenshein A.L., et al.,

Bacillus subtilis and other gram-positive bacteria. American society for Microbiology, Washington, p871(1993)).

Gene for CotE-antigen fusion protein is constructed by cloning the gene for surface antigen of HBV into pCotG-lacZ vector constructed in Example I-1. Thereafter, the constructed recombinant vector is transformed into Bacillus substilis and the transformants are cultured in GYS medium. And then, the antigen-displaying spores are purely isolated from culture medium by renografin gradients method.

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Example VIII: Protein Improvement Using Spore Displaying Protein of Interest

For example of application of the present invention to high-throughput screening of target protein and to protein improvement, GFP (Green Fluorescence Protein) was used as follows:

VIII-1: Construction of Vector for GFP Display on Spore Surface

gfp gene was cloned into pCSK-CotG vector constructed in Example IV-1 and the following sub-cloning procedures were performed for display on spore surface. Each primer was prepared for the purpose of fusing cotG gene to EGFP and GFPuv genes. The fluorescence intensity of EGFP (Excit./Emis. Maxima (nm): 488/509; Clontech, USA) has 35-fold stronger

than that of wild type GFP and thus results in detection even in FITC filter and GFPuv (Excit./Emis. Maxima (nm): 395/509; Clontech, USA) is detectable with UV. For further manipulation, NheI and HindIII restriction sites were inserted into primers for egfp gene (SEQ ID NOs:39 and 40) and PstI and EcoRI restriction sites were inserted in primers for gfpvu gene (SEQ ID NOs:41 and 42).

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Each of egfp (800 bp) and gfpuv (720 bp) genes was amplified by PCR (MJ Research PTC- 100^{TM} programmable Thermal Controller; 95% 30 sec, 55% 30 sec, 72% 2 min, 25 cycles) using Pfu Turbo polymerase (Stratagene, USA) and pEGFP-C1 (Clontedch, USA) or pGFPuv (Clontech, USA) as template.

Thereafter, pCSK-CotG-EGFP or pCSK-CotG-GFPuv vectors were constructed by cloning the restricted PCR products into NheI/HindIII (egfp gene) or PstI/EcoRI (gfpuv gene) restriction sites of pCSK-CotG vector.

VIII-2: Display and Confirmation of GFP on Spore Surface

The constructed vectors were transformed into Bacillus subtilis DB104 by natural transformation. Transformants were selected on LB agar plate containing 5 μ g/ml chloroamphenicol. Through the selection, Bacillus subtilis DB104-SDG-EGFP strain for display of EGFP and Bacillus subtilis DB104-SDG-GFPuv strain for display of GFPvu on spore surface were obtained. As control strains, Bacillus subtilis DB104-SDC

strain transformed with only pCSK vector and Bacillus subtilis DB104-SDG strain transformed for expressing only CotG protein were prepared.

For analysis of GFP display on spore surface, the above Bacillus subtilis DB104-SDC, -SDG, -SDG-EGFP and -SDG-GFPuv were inoculated into LB broth containing 5 $\mu g/ml$ chloroamphenicol and spores were then purified as described in Example V-4.

Subsequently, the display of GFP on spore surface was analyzed by measuring GFP fluorescence with flow cytometry in similar manner to Example IV (Fig. 16). In Fig. 16, curves (1)-(4) indicate the results of spores of DB104-SDC DB104-SDG, DB104-SDG-GFPuv and DB104-SDG-EFGP, respectively.

As shown in Fig. 16, the fluorescent intensity of spores derived from DB1047-SDG-EGFP (recombinant strain for EGFP-spore surface display) and DB104-SDG-GFPuv (recombinant strain for GFPuv-spore surface display) is significantly higher than that of DB104-SDC and DB104-SDG as control. In above results, the successful display of EGFP or GFPuv is validated by noticeable change of peaks indicating fluorescence in spore on its surface displaying EGFP or GFPuv comparing to controls.

VIII-3: Improvement of GFP

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For the purpose of GFP improvement, error prone PCR was

performed with template of pGFPuv vector (Clontech, USA) containing gfpuv gene using primers of SEQ ID NOs: 42 and 43. PCR mixture was prepared by mixing 0.3 μ M of each primers, 5 ng of DNA template, PCR solution (10mM Tris(pH 8.3), 50 mM KCl, 7 mM MgCl₂, 0.01% (w/v) gelatin), 0.2 mM dGTP, 0.2 mM dATP, 1 mM dTTP, 1 mM dCTP, 0.15 mM MnCl₂, 5 U Taq polymerase from Bioneer (Korea) and DW up to 100 μ L. Total 13 cycles of PCR was performed under condition of denaturation for 30 sec at 94°C, annealing for 30 sec at 50°C and extension for 1 min at 72°C.

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Subsequently, the gfpuv genes were discarded from pCSK-CotG-GFPuv vectors by restriction with PstI/EcoRI, the above PCR-amplified inserts were cloned into the vectors with the same restriction sites and Bacillus substilis DB104 was cloned by the natural transformed with vectors transformation to construct gfpuv library displayed on spore surface. Then, the prepared library was inoculated into GYS medium for sporulation and pure spores were isolated as described in Example V-4. Transformant spores displaying improved GFP variant measuring GFP were screened by fluorescence with flow cytometry (Figs. 17a to 17d). Figs. 17a to 17d indicates the analysis of flow cytometry from Bacillus subtilis DB104-SDC, DB104-SDG-GFPuv, DB104-SDG-EGFP and DB104-SDG-GFP with gfp library subject to error prone PCR, respectively.

To isolate spores with higher fluorescent intensity than spores derived from DB104-SDG-EGFP and DB104-SDG-GFP control strains, the isolation of spores with higher fluorescence (region R1) among spores displaying GFP library was repeated several times.

It is understood that using the above method, the improved GFP protein exhibiting higher fluorescence intensity or fluorescence with different wavelength can be screened in a high-throughput manner.

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Example VIII: Protein Array Using Spores Displaying on Its Surface Protein of Interest

106-109 spores displaying monoclonal antibodies against surface antigen of HBV are attached onto glass substrate for protein array (BMS, Germany) with aldehyde functional group on its surface using automated array apparatus. The attachment is made in a form of covalent linkage, which is Schiff base between amino group of protein on spore surface and aldehyde group on surface of slide glass. Although the displayed proteins attached on solid surface may be inactivated, they may have an orientation.

The protein array kit manufactured according to the present invention has a variety of applicable fields including diagnosis, analysis of gene expression, analysis of interaction between proteins, analysis of interaction

between protein and ligand, study on metabolism, screening novel or improved enzymes, combinatorial biochemical synthesis and biosensor.

Example IX: Production of Antibody Using Spores Displayig Antigen

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The spores on its surface displaying surface antigen of HBV isolated in Example VII are suspended in PBS and the same volume of complete Freund's adjuvant is added. Thereafter, the mixture is well agitated to make emulsion formulation and the emulsion is injected i.v. into BALB/c mice with age of 6-8 week. After 4 weeks of the injection, the secondary administration is performed. Then, the additional boosting injection is performed about 2-3 times for induction of antibody.

As described above, the display method on spore surface of the present invention provides improvements in: a resistance against physiochemical change in environment of display host, a diversity of displayable proteins, a viability of display host and rapidity of screening.

Throughout this application, various patents and publications are referenced and citations are provided in parentheses. The disclosure of these patents and

publications in their entities are hereby incorporated by references into this application in order to more fully describe this invention and the state of the art to which this invention pertains.

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Having described a preferred embodiment of the present invention, it is to be understood that variants and modifications thereof falling within the spirit of the invention may become apparent to those skilled in this art, and the scope of this invention is to be determined by appended claims and their equivalents.

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BUDAPEST TREATY ON THE INTERNATIONAL RECOGNITION OF THE DEPOSIT OF MICROORGANISMS FOR THE PURPOSE OF PATENT PROCEDURE

INTERNATIONAL FORM

RECEIPT IN THE CASE OF AN ORIGINAL DEPOSIT

issued pursuant to Rule 7.1

TO: PAN. Jae-Gu

=380-43. Doryong-dong, Yusong-ku, Taejon 305-340.

Republic of Korea

I. IDENTIFICATION OF THE MICROORGANISM

Identification reference given by the DEPOSITOR:

Bacillus subtilis GFSD18

Accession number given by the INTERNATIONAL DEPOSITARY AUTHORITY:

KCTC 0887BP

II. SCIENTIFIC DESCRIPTION AND/OR PROPOSED TAXONOMIC DESIGNATION

. The microorganism identified under I above was accompanied by:

[x] a scientific description

[] a proposed taxonomic designation

Mark with a cross where applicable)

III. RECEIPT AND ACCEPTANCE

This International Depositary Authority accepts the microorganism identified under I above, which was received by it on November 13 2000.

IN RECEIPT OF REQUEST FOR CONVERSION

The microorganism identified under I above was received by this International Depositary. Authority on and a request to convert the original deposit to a deposit under the Budapest Treaty was received by it on

V. INTERNATIONAL DEPOSITARY AUTHORITY

Name: Korean Collection for Type Cultures

Signature(s) of person(s) having the power to represent the International Depositary:
Authority of authorized official(s):

Address: Korea Research Institute of

Bioscience and Biotechnology (KRIBB)

=52. Oun-dong, Yusong-ku.

Taejon 305-333.
Republic of Korea

BAE. Kyung Sook. Director Date: November 16 2000

scie page

What is claimed is:

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1. A method for displaying a protein of interest on spore surface, which comprises the steps of:

- (i) preparing a vector for spore surface display comprising a gene construct containing a gene encoding spore coat protein and a gene encoding a protein of interest, wherein, when expressed, the gene construct expresses a fusion protein between the spore coat protein and the protein of interest;
- (ii) transforming a host cell with the vector for spore surface display;
- (iii) displaying the protein of interest on a surface of a spore of the host cell; and
- (iv) recovering the spore displaying on its surface the protein of interest.
- 2. A method for improving a protein of interest, which comprises the steps of:
 - (i) constructing a gene library of the protein of interest;
 - (ii) preparing a vector by linking the gene library to a gene encoding spore coat protein;
 - (iii) transforming a spore-forming host cell with the vector;
- 25 (iv) forming a spore in the transformed host cell and

displaying the protein of interest on a surface of the spore;

- (v) recovering the spore displaying on its surface the protein of interest; and
- (vi) screening the spore displaying a variant of the protein of interest having a desired property.

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- 3. The method according to claim 2, wherein the screening is performed by means of measuring an activity of the protein or flow cytometry.
- 4. A method for improving a protein of interest using a resistance property of spore, which comprises the steps of:
 - (i) constructing a gene library of the protein of interest;
 - (ii) preparing a vector by linking the gene library to a gene encoding spore coat protein;
 - (iii) transforming a spore-forming host cell with the vector;
- (iv) forming a spore in the transformed host cell and displaying the protein of interest on a surface of the spore;
 - (v) treating the spore displaying on its surface the protein of interest with one or more selected from the group consisting of organic solvent, heat, acid, base,

oxidant, dryness, surfactant and protease;

(vi) recovering the spore displaying on its surface the protein of interest; and

(vii) screening the spore displaying a variant of the protein of interest having a resistance to the treatment.

5. The method according to claim 4, wherein the screening is performed using an activity of the protein or a structural stability of the protein.

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- 6. A method for bioconversion, which comprises the steps of:
 - (i) preparing a vector for spore surface display comprising a gene construct containing a gene encoding spore coat protein and a gene encoding a protein of interest conducting a bioconversion reaction, wherein, when expressed, the gene construct expresses a fusion protein between the spore coat protein and the protein of interest;
 - (ii) transforming a host cell with the vector for spore surface display;
 - (iii) displaying the protein of interest on a surface of a spore of the host cell;
 - (iv) recovering the spore displaying on its surface the protein of interest; and
- (v) performing the bioconversion reaction using the spore

displaying on its surface the protein of interest.

7. A method for preparing protein microarray, which comprises the steps of:

(i) preparing a vector for spore surface display comprising a gene construct containing a gene encoding spore coat protein and a gene encoding antibody or antigen having binding affinity to a protein to be analyzed, wherein, when expressed, the gene construct expresses a fusion protein between the spore coat protein

and the antibody or antigen;

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- (ii) transforming a host cell with the vector for spore surface display;
- (iii) displaying the antibody or antigen on a surface of a spore of the host cell;
- (iv) recovering the spore displaying on its surface the antibody or antigen; and
- (v) immobilizing onto a solid surface the spore displaying on its surface the antibody or antigen.
- 8. A method producing an antibody to antigen in vertebrates, which comprises the steps of:
- (i) preparing a vector for spore surface display comprising a gene construct containing a gene encoding spore coat protein and a gene encoding the antigen,

wherein, when expressed, the gene construct expresses a fusion protein between the spore coat protein and the antigen;

(ii) transforming a host cell with the vector for spore surface display;

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- (iii) displaying the antigen on a surface of a spore of the host cell;
- (iv) recovering the spore displaying on its surface the antigen; and
- (v) administering to vertebrates a composition containing an immunologically effective amount of the spore displaying on its surface the antigen.
- 9. A method for preparing a whole cell absorber, which comprises the steps of:
 - (i) preparing a vector for spore surface display comprising a gene construct containing a gene encoding spore coat protein and a gene encoding a protein having a binding affinity to a certain substance, wherein, when expressed, the gene construct expresses a fusion protein between the spore coat protein and the protein;
 - (ii) transforming a host cell with the vector for spore surface display;
- (iii) displaying the protein on a surface of a spore of the host cell;

(iv) recovering the spore displaying on its surface the protein; and

(v) immobilizing onto a carrier the spore displaying on its surface the protein.

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- 10. The method according to any one of claims 1-9, wherein the gene encoding spore coat protein is derived from a spore-forming Gram negative bacterium including Myxococcus, a spore-forming Gram positive bacterium including Bacillus, a spore-forming Actionmycete, a spore-forming yeast or a spore-forming fungus.
- 11. The method according to claim 10, wherein the gene encoding spore coat protein is derived from a spore-forming Gram positive bacterium.
- 12. The method according to claim 11, wherein the gene encoding spore coat protein is derived from Bacillus.
- 20 13. The method according to any one of claims 1-9, wherein the gene encoding spore coat protein is selected from the group consisting of cotA, cotB, cotC, cotD, cotE, cotF, cotG, cotH, cotJA, cotJC, cotK, cotL, cotM, cotS, cotT, cotV, cotW, cotX, cotY, cotZ, spoIVA, spoVID and sodA.

14. The method according to any one of claims 1-9, wherein the gene encoding spore coat protein is selected from the group consisting of cotA, cotE, cotF, cotG, cotH, cotJA, cotJC, cotK, cotL, cotM, cotS, cotT, cotV, cotW, cotX, cotY, cotZ, spoIVA, spoVID and sodA.

15. The method according to any one of claims 1-9, wherein the gene encoding spore coat protein is a modified form or a recombinant form of one selected from the group consisting of cotA, cotB, cotC, cotD, cotE, cotF, cotG, cotH, cotJA, cotJC, cotK, cotL, cotM, cotS, cotT, cotV, cotW, cotX, cotY, cotZ, spoIVA, spoVID and sodA, in which the modified form or the recombinant form has a more compatibility for spore surface display relative to wild type genes.

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16. The method according to claim 15, wherein the modified form of the gene encoding spore coat protein is obtained by a method selected from the group consisting of DNA shuffling method, StEP method, RPR method, molecular breeding method, ITCHY method, error-prone PCR, point mutagenesis, nucleotide mutagenesis, combinatorial cassette mutagenesis and other suitable random mutagenesis.

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17. The method according to any one of claims 1-9, wherein the gene encoding spore coat protein is selected from the

group consisting of cotA, cotB, cotC, cotD, cotE, cotF, cotG, cotH, cotJA, cotJC, cotK, cotL, cotM, cotS, cotT, cotV, cotW, cotX, cotY, cotZ, spoIVA, spoVID and sodA, in which the gene has a substituted promoter for its promoter to enhance spore surface display relative to wild type genes.

- 18. The method according to claim 13, wherein the gene encoding spore coat protein is cotE or cotG.
- 10 19. The method according to claim 14, wherein the gene encoding spore coat protein is cotE or cotG.
 - 20. The method according to claim 15, wherein the gene encoding spore coat protein is cotE or cotG.

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- 21. The method according to claim 16, wherein the gene encoding spore coat protein is cotE or cotG.
- 22. The method according to claim 16, wherein the gene encoding spore coat protein is cotE or cotG.
 - 23. The method according to any one of claims 1-5, wherein the protein of interest is selected from the group consisting of enzyme, enzyme inhibitor, hormone, hormone analogue, hormone receptor, signal transduction protein,

antibody, monoclonal antibody, antigen, adhesive protein, structural protein, regulatory protein, toxin protein, cytokine, transcription regulatory protein, blood clotting protein, plant protection-inducing protein and fragments thereof.

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- 24. The method according to any one of claims 1-9, wherein the host cell is selected from the group consisting of a spore-forming Gram negative bacterium including Myxococcus, a spore-forming Gram positive bacterium including Bacillus, a spore-forming Actionmycete, a spore-forming yeast or a spore-forming fungus.
- 25. The method according to claim 24, wherein the host cell is a spore-forming Gram positive bacterium.
 - 26. The method according to claim 25, wherein the host cell is Bacillus.
- 27. The method according to any one of claims 1-9, wherein the spore is reproductive or non-reproductive one.
 - 28. The method according to any one of claims 1-9, wherein the recovering is performed in such a manner that the display of the protein of interest on the spore surface is

maximized by regulating culture time, after which culturing is terminated and the spore is then recovered.

- 29. The method according to any one of claims 2-5, wherein library is performed by a constructing a gene the 5 mutagenesis of the gene encoding the protein of interest of wild type, in which the mutagenesis is selected from the group consisting of DNA shuffling method, StEP method, RPR method, molecular breeding method, ITCHY method, error-prone mutagenesis, nucleotide mutagenesis, point PCR, 10 combinatorial cassette mutagenesis and other suitable random mutagenesis.
- protein of interest, characterized in that the transformant is produced by transformation with a vector for spore surface display containing (i) a gene encoding a protein of interest and (ii) a gene encoding spore coat protein is selected from the group consisting of cotA, cotB, cotC, cotD, cotE, cotF, cotG, cotH, cotJA, cotJC, cotK, cotL, cotM, cotS, cotT, cotV, cotW, cotX, cotY, cotZ, spoIVA, spoVID and sodA, in which when expressed, a fusion protein between the spore coat protein and the protein of interest is expressed.
 - 31. The transformant according to claim 30, wherein the

transformant is derived from a variant mutated to enhance spore surface display.

- 32. The transformant according to claim 31, wherein the mutation to enhance spore surface display eliminates a production of extracellular secretory protease in the transformant, so that the protein of interest displayed on spore surface is stably maintained.
- 33. The transformant according to claim 31, wherein the mutation to enhance spore surface display eliminates a production of intracellular protease in the transformant.
- 34. A spore for spore surface display of a protein of interest, characterized in that the spore displays the protein of interest on its surface.
 - 35. The spore according to claim 34, wherein the spore is reproductive or non-reproductive one.
 - 36. The spore according to claim 35, wherein the spore is non-reproductive one by virtue of one or more methods selected from the group consisting of genetic method, chemical method and physical method.

37. The spore according to claim 36, wherein the genetic method to make the spore non-reproductive is accomplished by deleting a gene involved in reproduction of spore.

- 38. The spore according to claim 34, wherein the spore is derived from a variant mutated to increase its agglutination property.
- 39. The spore according to claim 38, wherein the increase of the agglutination property in the spore is accomplished by one or more methods selected from the group consisting of genetic method, chemical method and physical method.
- 40. A vector for spore surface display, characterized in
 that the vector comprises a replication origin, an
 antibiotic-resistance gene, a restriction site, a gene
 encoding a spore coat protein, a gene encoding a protein of
 interest and a promoter operatively linked to the gene
 encoding a spore coat protein, in which when expressed, a

 20 fusion protein between the spore coat protein and the
 protein of interest is expressed.
 - 41. The vector according to claim 40, wherein the gene encoding a spore coat protein is selected from the group consisting of cotA, cotB, cotC, cotD, cotE, cotF, cotG, cotH,

cotJA, cotJC, cotK, cotL, cotM, cotS, cotT, cotV, cotW, cotX, cotY, cotZ, spoIVA, spoVID and sodA.

42. The vector according to claim 41, wherein the gene encoding a spore coat protein is cotE or cotG.

1 / 15 FIG. 1

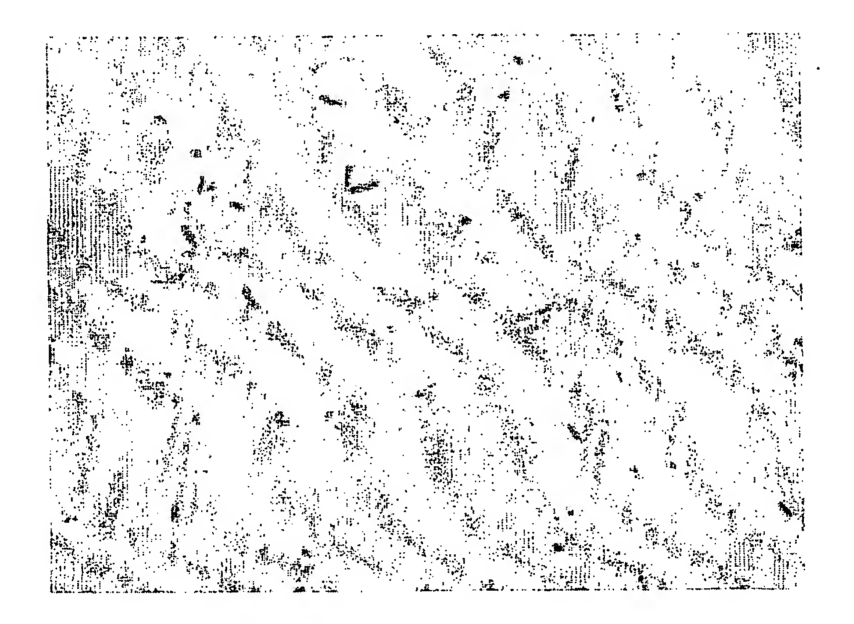
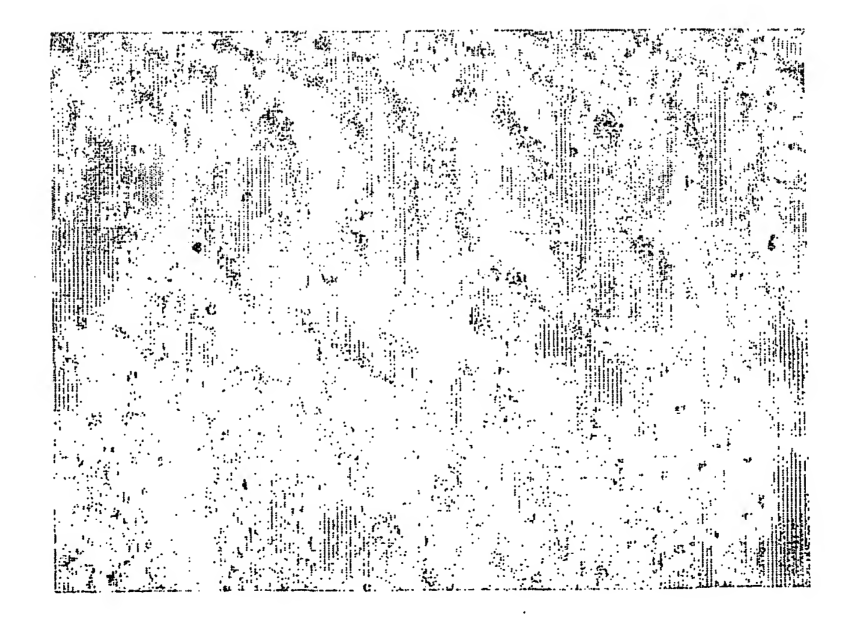
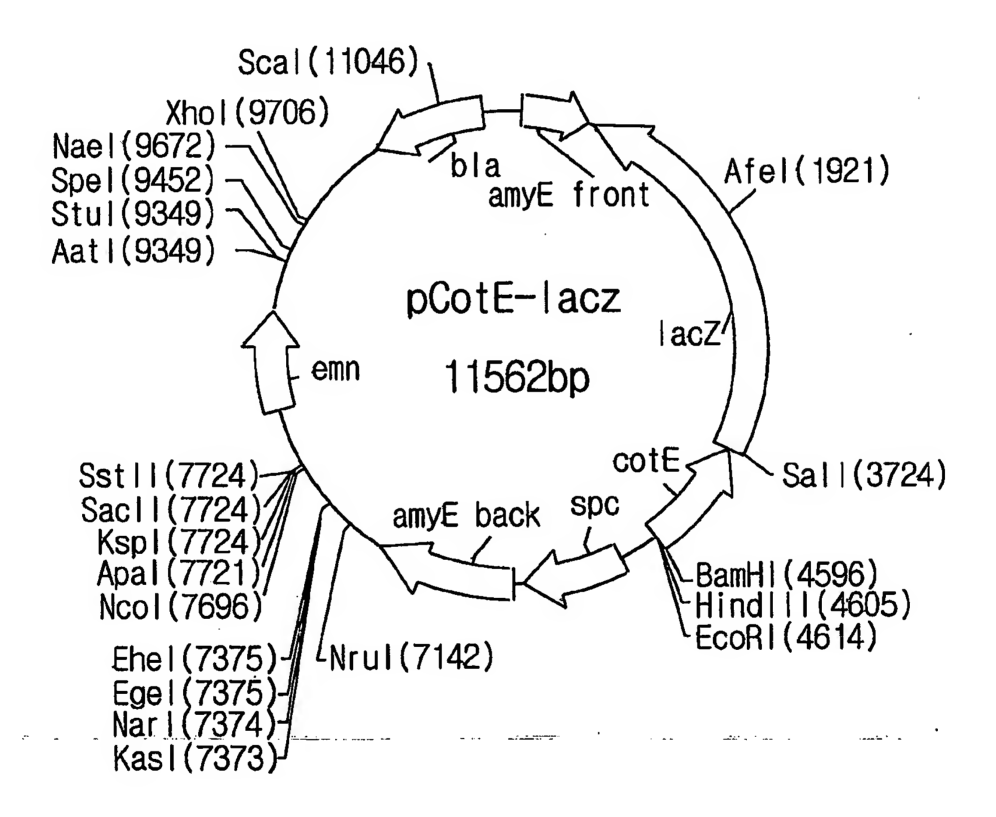


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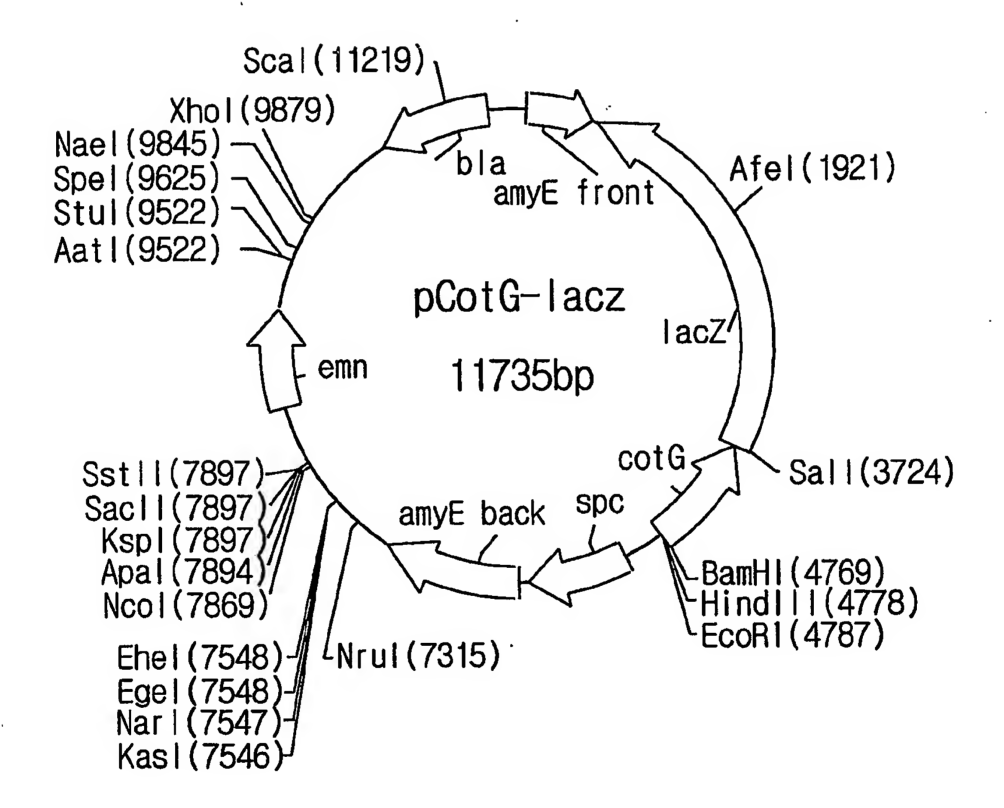


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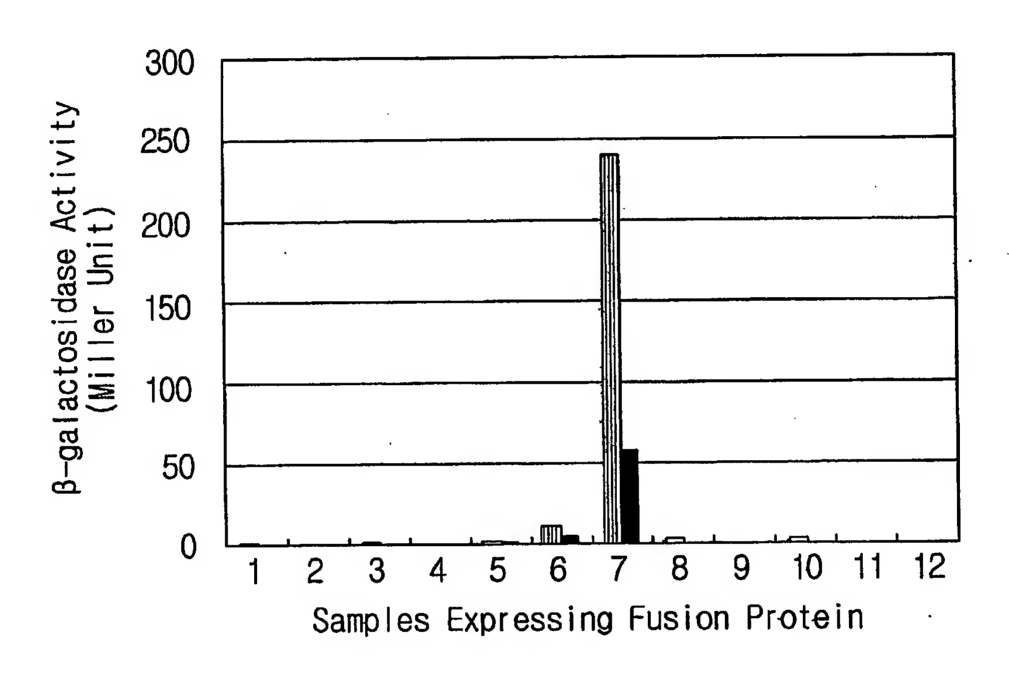
2 / 15 FIG.3



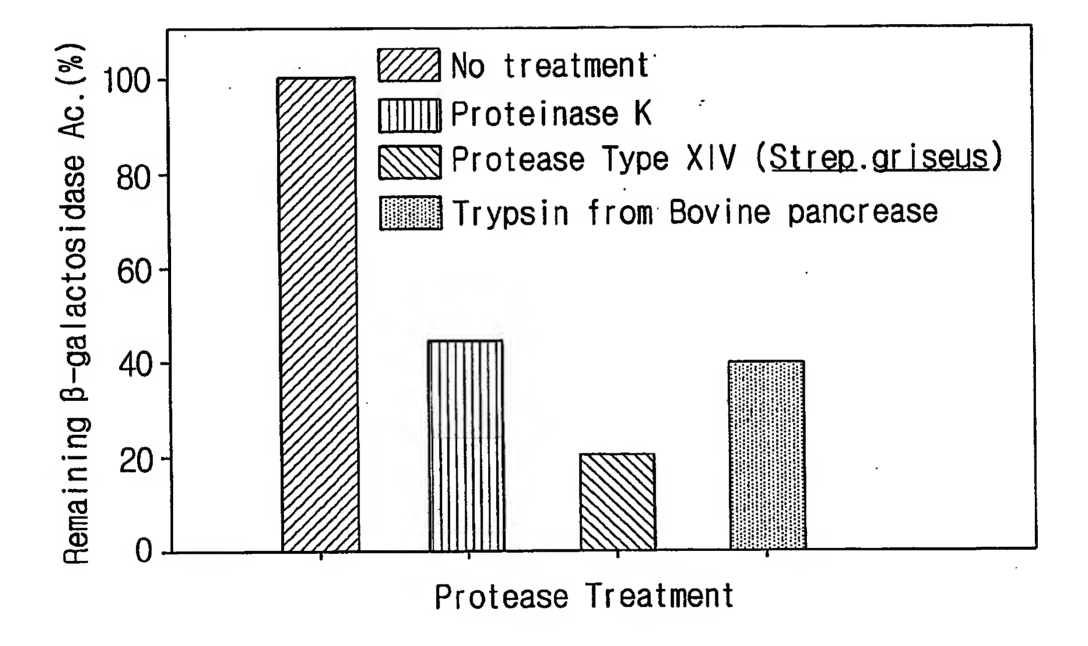
3 / 15 FIG.4



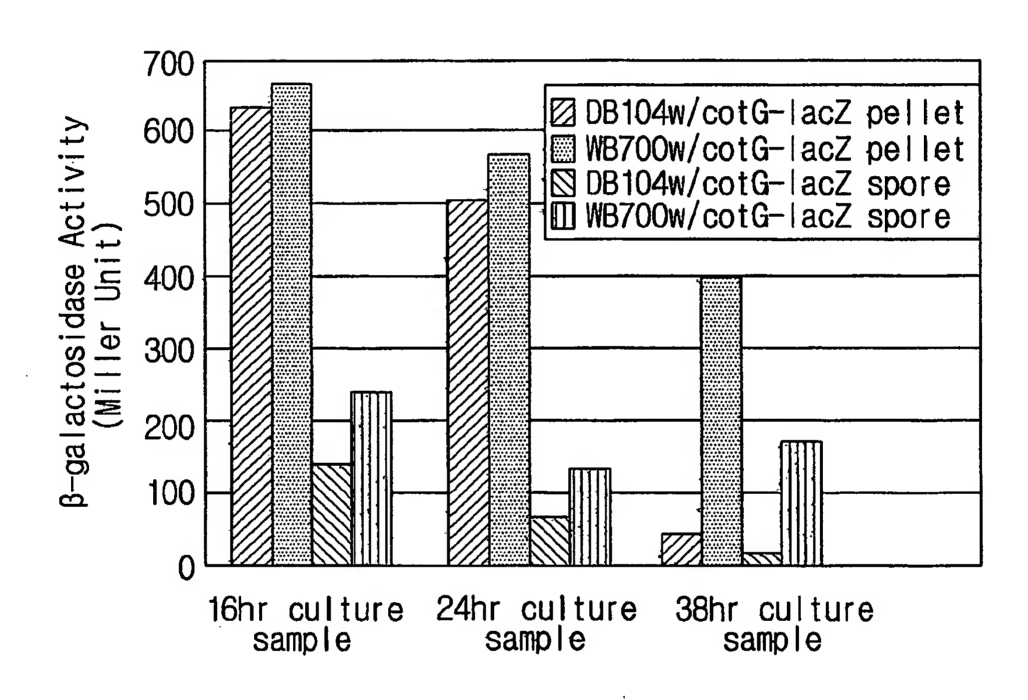
4 / 15 FIG.5



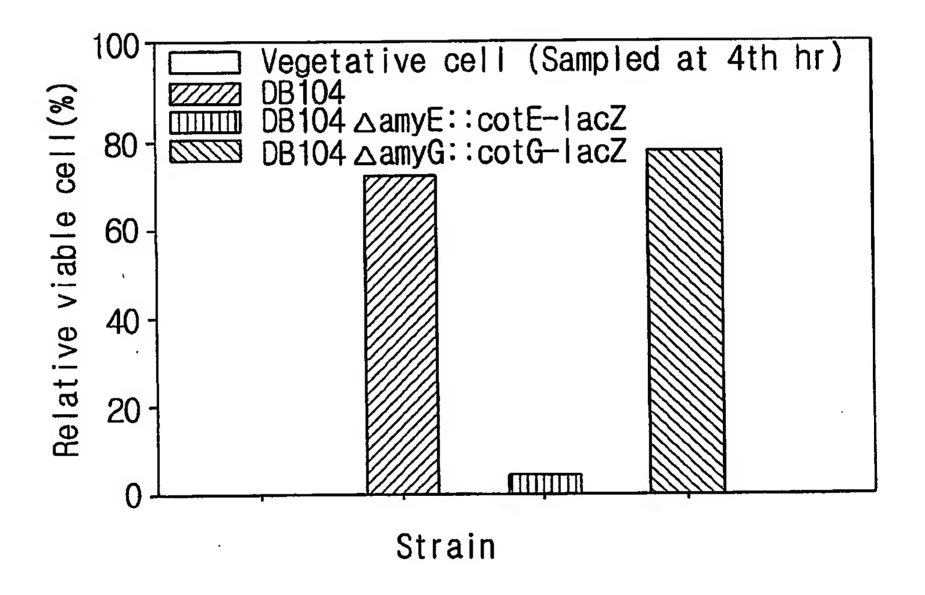
5 / 15 FIG.6



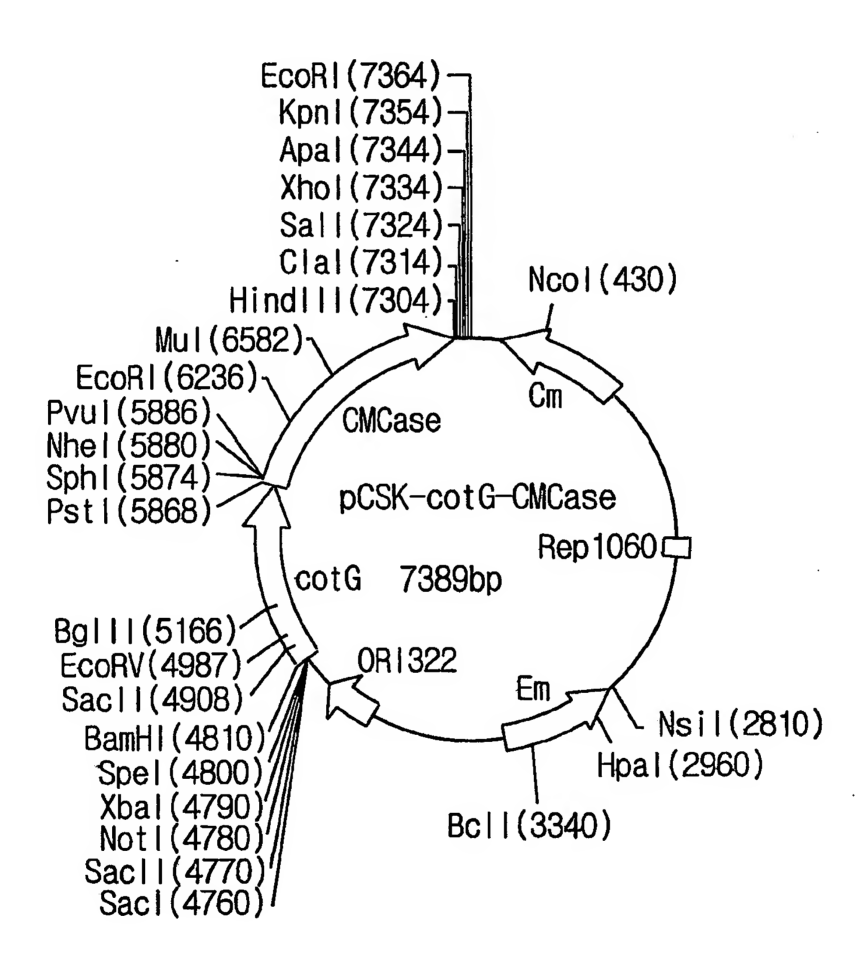
6 / 15 FIG. 7



7 / 15 FIG.8



8 / 15 FIG.9



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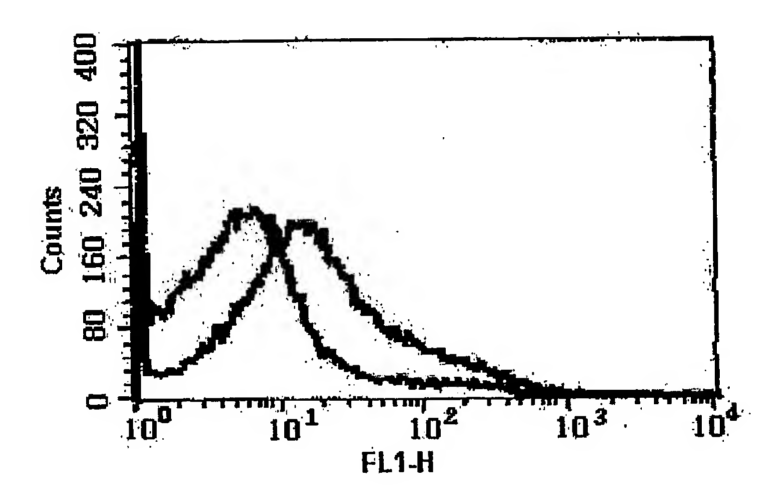
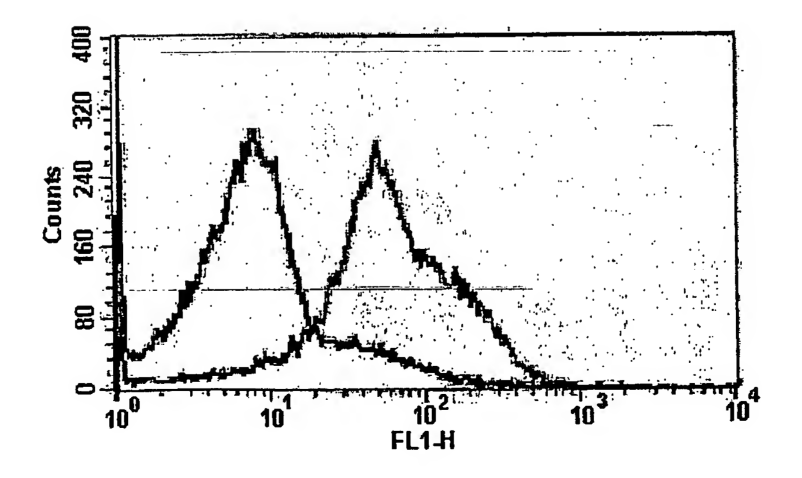
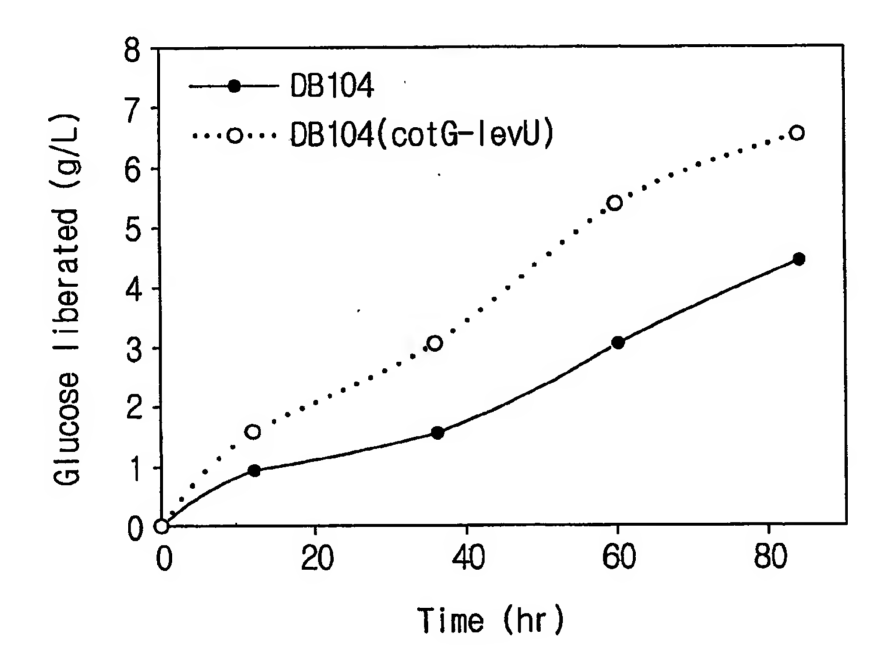


FIG.11



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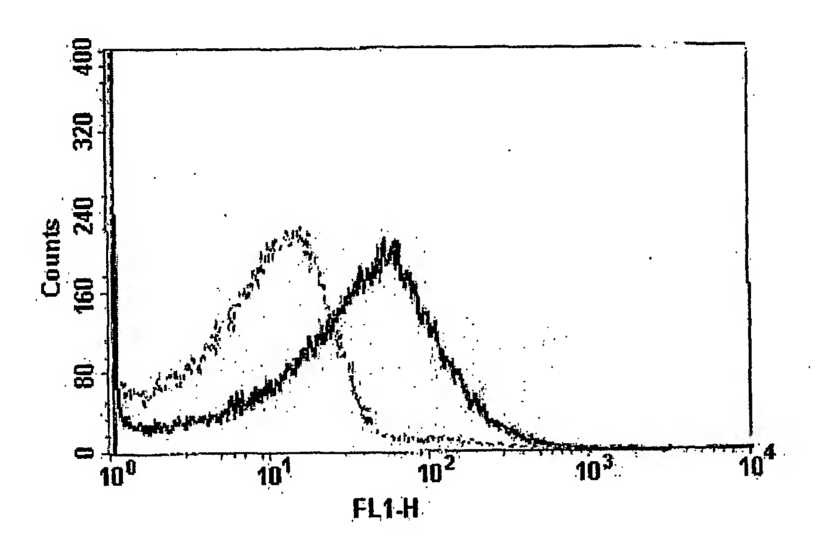
10 / 15 FIG. 12



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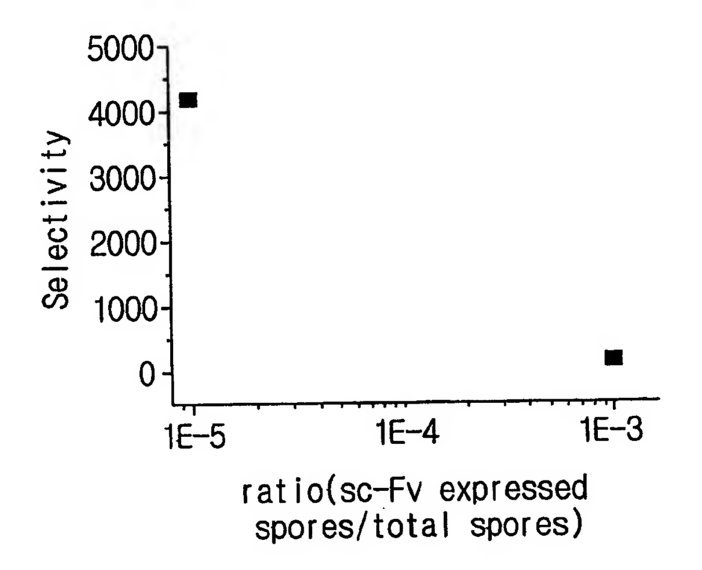
11 / 15 FIG. 13

PCT/KR01/02124



PCT/KR01/02124

12 / 15 FIG. 14



<u>.</u>

13 / 15 FIG. 15

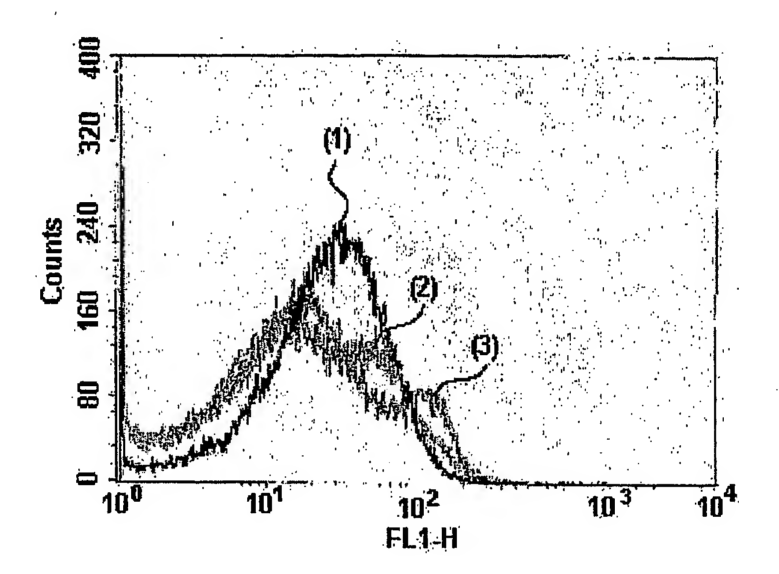
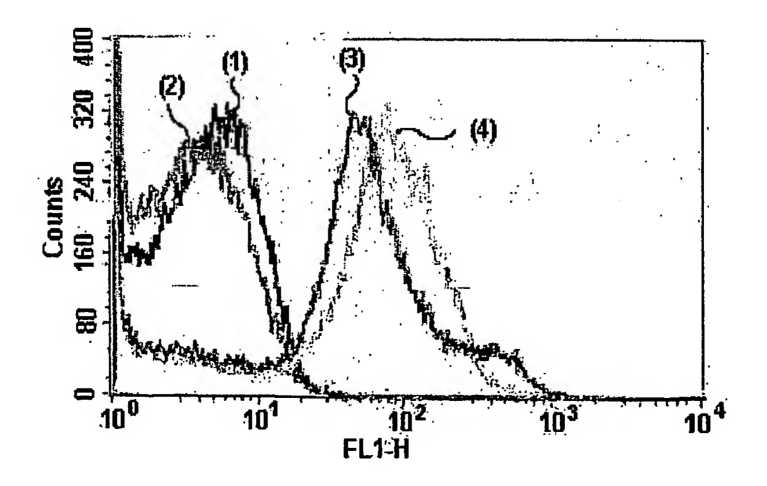


FIG.16



PCT/KR01/02124

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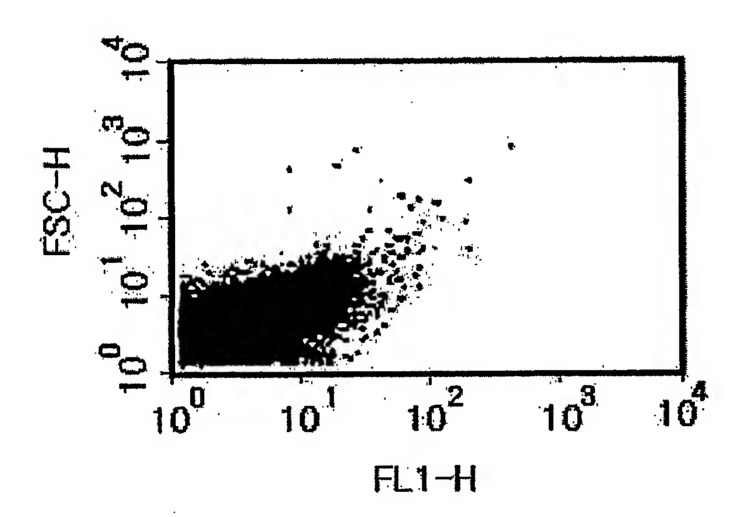
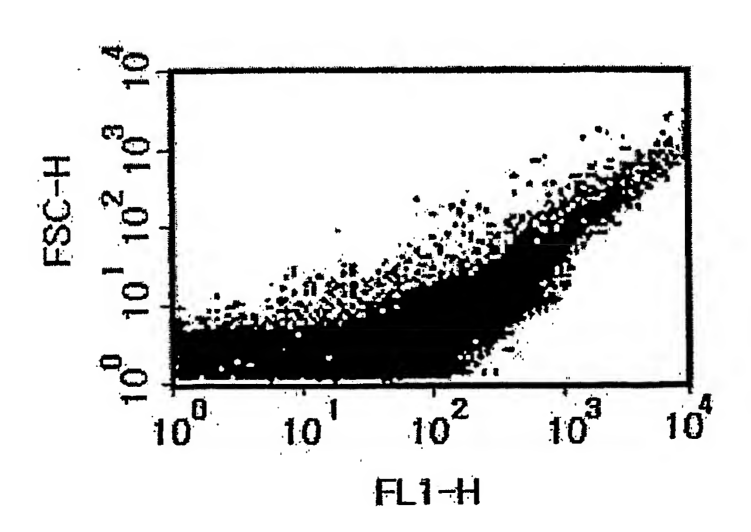


FIG.17b



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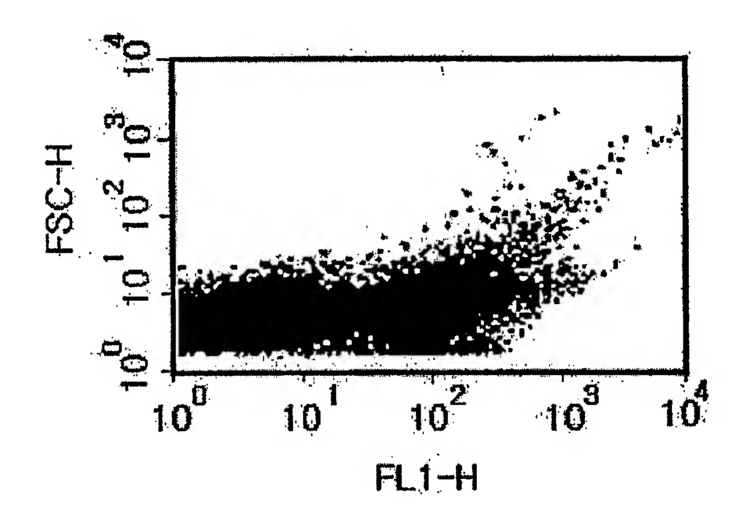
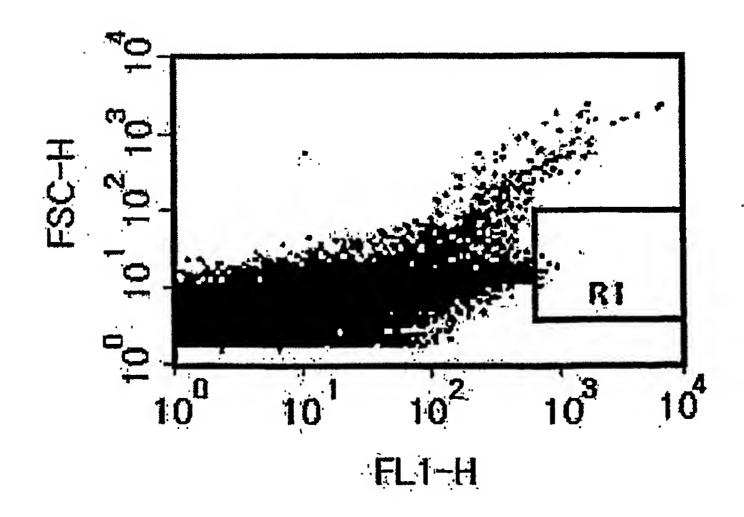


FIG.17d



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gcg ccc atc tac				•	1169
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Val Asp Glu Ser 300		aly Gln Thr Arg Ile 305	e lle Phe Asp 310	
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010	020	•	020	
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330	335		10	
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ttc cgt gac gtc to Phe Arg Asp Val			_	1505
380		385	390	
ttc cat gtt gcc ac Phe His Val Ala 1 395	ct cgc ttt aat gat Thr Arg Phe Asn A 400	Asp Asp Phe Ser		1553 J
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	g cag ggt gaa ac			1649
425	rp Gln Gly Glu Tl 430	nr Gin vai Ala Se 435	i Giy ini Ala	440
TLU	1 00	400		770
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Pro Phe Gly Gly Glu 44	·	rg Gly Gly Tyr 450	Ala Asp Arg 455	
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atc ccg aat ctc tat o				1793
475	480	ia Loa i no i ni	485	
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etg tge gac egc tac Leu Cys Asp Arg Ty				2081
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Thr His Gly Met Val Pro Met Asn Arg Leu Thr Asp Asp Pro Arg Trp cta ccg gcg atg agc gaa cgc gta acg cga atg gtg cag cgc gat cgt Leu Pro Ala Met Ser Glu Arg Val Thr Arg Met Val Gln Arg Asp Arg aat cac ccg agt gtg atc atc tgg tcg ctg ggg aat gaa tca ggc cac Asn His Pro Ser Val IIe IIe Trp Ser Leu Gly Asn Glu Ser Gly His ggc gct aat cac gac gcg ctg tat cgc tgg atc aaa tct gtc gat cct Gly Ala Asn His Asp Ala Leu Tyr Arg Trp Ile Lys Ser Val Asp Pro tee ege eeg gtg eag tat gaa gge gge gga gee gae ace acg gee ace Ser Arg Pro Val Gln Tyr Glu Gly Gly Gly Ala Asp Thr Thr Ala Thr gat att att tgc ccg atg tac gcg cgc gtg gat gaa gac cag ccc ttc Asp lle lle Cys Pro Met Tyr Ala Arg Val Asp Glu Asp Gln Pro Phe ccg gct gtg ccg aaa tgg tcc atc aaa aaa tgg ctt tcg cta cct gga Pro Ala Val Pro Lys Trp Ser lle Lys Lys Trp Leu Ser Leu Pro Gly gag acg cgc ccg ctg atc ctt tgc gaa tac gcc cac gcg atg ggt aac Glu Thr Arg Pro Leu lie Leu Cys Glu Tyr Ala His Ala Met Gly Asn agt ctt ggc ggt ttc gct aaa tac tgg cag gcg ttt cgt cag tat ccc Ser Leu Gly Gly Phe Ala Lys Tyr Trp Gln Ala Phe Arg Gln Tyr Pro cgt tta cag ggc ggc ttc gtc tgg gac tgg gtg gat cag tcg ctg att

7,

Arg Leu Gin Gly Gly Phe Val Trp Asp Trp Val Asp Gin Ser Leu lie aaa tat gat gaa aac ggc aac ccg tgg tcg gct tac ggc ggt gat ttt Lys Tyr Asp Glu Asn Gly Asn Pro Trp Ser Ala Tyr Gly Gly Asp Phe ggc gat acg ccg aac gat cgc cag ttc tgt atg aac ggt ctg gtc ttt Gly Asp Thr Pro Asn Asp Arg Gln Phe Cys Met Asn Gly Leu Val Phe gcc gac cgc acg ccg cat cca gcg ctg acg gaa gca aaa cac cag cag Ala Asp Arg Thr Pro His Pro Ala Leu Thr Glu Ala Lys His Gln Gln cag ttt ttc cag ttc cgt tta tcc ggg caa acc atc gaa gtg acc agc Gln Phe Phe Gln Phe Arg Leu Ser Gly Gln Thr Ile Glu Val Thr Ser gaa tac ctg ttc cgt cat agc gat aac gag ctc ctg cac tgg atg gtg Glu Tyr Leu Phe Arg His Ser Asp Asn Glu Leu Leu His Trp Met Val gcg ctg gat ggt aag ccg ctg gca agc ggt gaa gtg cct ctg gat gtc Ala Leu Asp Gly Lys Pro Leu Ala Ser Gly Glu Val Pro Leu Asp Val gct cca caa ggt aaa cag-ttg att-gaa-ctg cct gaa cta ccg cag ccg Ala Pro Gin Gly Lys Gin Leu Ile Giu Leu Pro Giu Leu Pro Gin Pro gag age gee ggg caa etc tgg-etc aca gta-ege gta gtg caa eeg aac Glu Ser Ala Gly Gln Leu Trp Leu Thr Val Arg Val Val Gln Pro Asn

gcg acc gca tgg tca gaa gcc ggg cac atc agc gcc tgg cag cag tgg

Ala Thr Ala Trp Ser Glu Ala Gly His Ile Ser Ala Trp Gln Gln Trp .875 cgt ctg gcg gaa aac ctc agt gtg acg ctc ccc gcc gcg tcc cac gcc Arg Leu Ala Glu Asn Leu Ser Val Thr Leu Pro Ala Ala Ser His Ala atc ccg cat ctg acc acc agc gaa atg gat ttt tgc atc gag ctg ggt lle Pro His Leu Thr Thr Ser Glu Met Asp Phe Cys lle Glu Leu Gly aat aag cgt tgg caa ttt aac cgc cag tca ggc ttt ctt tca cag atg Asn Lys Arg Trp Gln Phe Asn Arg Gln Ser Gly Phe Leu Ser Gln Met tgg att ggc gat aaa aaa caa ctg ctg acg ccg ctg cgc gat cag ttc Trp lle Gly Asp Lys Lys Gln Leu Leu Thr Pro Leu Arg Asp Gln Phe acc cgt gca ccg ctg gat aac gac att ggc gta agt gaa gcg acc cgc Thr Arg Ala Pro Leu Asp Asn Asp Ile Gly Val Ser Glu Ala Thr Arg att gac cct aac gcc tgg gtc gaa cgc tgg aag gcg gcg ggc cat tac lle Asp Pro Asn Ala Trp Val Glu Arg Trp Lys Ala Ala Gly His Tyr cag gcc gaa gca gcg ttg ttg cag tgc acg gca gat aca ctt gct gat Gln Ala Glu Ala Ala Leu Leu Gln Cys Thr Ala Asp Thr Leu Ala Asp gcg gtg ctg att acg acc gct cac gcg tgg cag cat cag ggg aaa acc Ala Val Leu lle Thr Thr Ala His Ala Trp Gln His Gln Gly Lys Thr tta ttt atc agc cgg aaa acc tac cgg att gat ggt agt ggt caa atg

Leu Phe lle	Ser Arg Lys Thr 1020	Tyr Arg lle Asp 1025		n Met 030	
•	gtt gat gtt gaa (/al Asp Val Glu \ 5			o Ala	3473
cgg att ggo	ctg aac tgc car Leu Asn Cys Glr	g ctg gcg cag			3521
	tta ggg ccg ca Leu Gly Pro Gli 1070				3569 . ·
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Trp Tyr Ser Tyr Ala Asp Asn Thr Lys Thr Glu Val Val Thr Glu Arg
65 70 75 80

Val Lys Tyr Val Asp Val Ile Lys Leu Arg Tyr Arg Asp Asn Asn Tyr
85 90 95

55

50

60

Leu Asp Asp Glu His Glu Val Ile Ala Lys Val Leu Gln Gln Pro Asn Cys Leu Glu Val Thr lle Ser Pro Asn Gly Asn Lys lle Val Val Gln Ala Glu Arg Glu Phe Leu Ala Glu Val Val Gly Glu Thr Lys Val Val Val Glu Val Asn Pro Asp Trp Glu Glu Asp Asp Glu Glu Asp Trp Glu Asp Glu Leu Asp Glu Glu Leu Glu Asp lle Asn Pro Glu Phe Leu Val Gly Asp Pro Glu Glu Val Asp Arg Glu Asn Pro Gly Val Thr Gln Leu Asn Arg Leu Ala Ala His Pro Pro Phe Ala Ser Trp Arg Asn Ser Glu Glu Ala Arg Thr Asp Arg Pro Ser Gln Gln Leu Arg Ser Leu Asn Gly Glu Trp Arg Phe Ala Trp Phe Pro Ala Pro Glu Ala Val Pro Glu Ser Trp Leu Glu Cys Asp Leu Pro Glu Ala Asp Thr Val Val Val Pro Ser Asn Trp Gln Met His Gly Tyr Asp Ala Pro Ile Tyr Thr Asn Val Thr Tyr Pro lle Thr Val Asn Pro Pro Phe Val Pro Thr Glu Asn Pro Thr

Gly Cys Tyr	Ser Leu Thr Phe A	sn Val Asp Glu S	Ser Trp Leu Gin Glu	
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Gly Gln Thr	Arg lle lle Phe Asp	Gly Val Asn Ser	Ala Phe His Leu	
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Ser Glu Phe	Asp Leu Ser Ala f	Phe Leu Arg Ala	Gly Glu Asn Arg Le	u
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Ala Val Met	Val Leu Arg Trp Se	er Asp Gly Ser Ty	yr Leu Glu Asp Gln	
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Asp Met Tro	Ara Met Ser Glv I	le Phe Ara Aso \	/al Ser Leu Leu His	
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070	070	,		
Lve Pro Thr	Thr Gln lle Ser As	n Phe His Val Als	a Thr Aro Phe Asn	
385	390		95	400
000	030			400
Asp Asp Ph	a Sar Ara Ala Val	ou Glu Ala Glu I	Val Gln Met Cys Gly	,
Yah yah Lii	405	410	vai diir iviet dys dij 415	
	405	410	415	,
Challan Ara	An Turkey	Val The Val Car I	on Ten Cla Cla Cla	
Giu Leu Arg			eu Trp Gln Gly Glu	
	420	425	430	
			o o o o o o o o o o o o o o o o o o o	
	Ala Ser Gly Thr Al -			
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		_	rg Leu Asn Val Glu	
450	45	5	460	
Asn Pro Lys	s Leu Trp Ser Ala	Glu lle Pro Asn L	eu Tyr Arg Ala Val	
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Asn Gly Lys 515	Pro Leu Leu lle	Arg Gly Val Asr 520	n Arg His Glu H 525	is His
Pro Leu His 530	Gly Gln Val Met 5	Asp Glu Gln Tł 35	nr Met Val Gln A 540	asp lle
Leu Leu Met	Lys Gln Asn As	n Phe Asn Ala	Val Arg Cys Se	r His Tyr
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Pro Asn His	Pro Leu Trp Tyr	Thr Leu Cys A	sp Arg Tyr Gly	Leu Tyr
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Thr Arg Met	Val Gln Arg Asp	Arg Asn His P	ro Ser Val Ile III	e Trp
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Ser Leu-Gly	Asn Glu Ser Gly	y His Gly Ala As	sn His Asp Ala	Leu Tyr
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Arg Trp Ile l	_ys Ser Val Asp	Pro Ser Arg Pro	val Gln Tyr Gl	u Gly
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Trp Ser Ala 755		he Gly Asp Thr 760	Pro Asn Asp Arg 765	Gin
Phe Cys Me	t Asn Gly Leu Val 775		rg Thr Pro His Pro 780	Ala
Leu Thr Glu 785	Ala Lys His Gln G 790		e Gln Phe Arg Leu 795	Ser 800
Gly Gln Thr	lle Glu Val Thr Se 805	r Glu Tyr Leu P 810	he Arg His Ser As	p 815
Asn Glu Leu	Leu His Trp Met 820	Val Ala Leu As 825	p Gly Lys Pro Leu 830	Ala
Ser Gly Glu 835		Val Ala Pro Gln 840	Gly Lys Gln Leu I 845	le
Glu Leu Pro 850	Glu Leu Pro Gln 85		a Gly Gln Leu Trp 860	Leu

Thr Val Arg Val Val Gln Pro Asn Ala Thr Ala Trp Ser Glu Ala Gly His lie Ser Ala Trp Gln Gln Trp Arg Leu Ala Glu Asn Leu Ser Val Thr Leu Pro Ala Ala Ser His Ala lle Pro His Leu Thr Thr Ser Glu Met Asp Phe Cys Ile Glu Leu Gly Asn Lys Arg Trp Gln Phe Asn Arg Gin Ser Gly Phe Leu Ser Gin Met Trp lie Gly Asp Lys Lys Gin Leu Leu Thr Pro Leu Arg Asp Gln Phe Thr Arg Ala Pro Leu Asp Asn Asp lle Gly Val Ser Glu Ala Thr Arg lle Asp Pro Asn Ala Trp Val Glu Arg Trp Lys Ala Ala Gly His Tyr Gln Ala Glu Ala Ala Leu Leu Gln Cys Thr Ala Asp Thr Leu Ala Asp Ala Val Leu lle Thr Thr Ala His Ala Trp-Gln-His Gln Gly Lys Thr-Leu Phe Ile Ser Arg Lys Thr Tyr Arg lie Asp Gly Ser Gly Gln Met Ala lle Thr Val Asp Val Glu Val Ala Ser Asp Thr Pro His Pro Ala Arg Ile Gly Leu Asn Cys Gln Leu

Ala Gin Val Ala Giu Arg Val Asn Trp Leu Gly Leu Gly Pro Gin Giu 1060 1065 1070

Asn Tyr Pro Asp Arg Leu Thr Ala Ala Cys Phe Asp Arg Trp Asp Leu 1075 1080 1085

Pro Leu Ser Asp Met Tyr Thr Pro Tyr Val Phe Pro Ser Glu Asn Gly 1090 1095 1100

Leu Arg Cys Gly Thr Arg Glu Leu Asn Tyr Gly Pro His Gln Trp Arg 1105 1110 1115 1120

Gly Asp Phe Gln Phe Asn Ile Ser Arg Tyr Ser Gln Gln Gln Leu Met 1125 1130 1135

Glu Thr Ser His Arg His Leu Leu His Ala Glu Glu Gly Thr Trp Leu 1140 1145 1150

Asn Ile Asp Gly Phe His Met Gly Ile Gly Gly Asp Asp Ser Trp Ser 1155 1160 1165

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	at tta tac caa gag					571
Lys Asp 1	yr Leu Tyr Gln G 25	30	y Lys Lys .	Arg Ser His	Lys	
Lys Ser H	ac cgc act cac a is Arg Thr His Ly 40					619
Cys Ser H	c aaa aaa tot og Iis Lys Lys Ser A	rg Ser His Ly				667
55		60		65		

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Lys Lys Ser /	Arg Ser His Lys Ly	s Ser Tyr Cys	Ser His Lys Ly	s Ser	
70	75		80	85	
cgc agc cac	aaa aaa tog tac	ogt tot cac aa	a aaa tot ogo a	gc tat	763
Arg Ser His l	ys Lys Ser Tyr Ar	g Ser His Lys	Lys Ser Arg Se	r Tyr	
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tot t	on not tot ton one			101	011
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Lys Lys Ser	Tyr Arg Ser Tyr Ly				
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tac cat tot te	nc aaa aaa tot ogo	n ann tan aaa	aga tot tac tat	tot	859
	Tyr Lys Lys Ser A				000
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	Lys Ser Arg Ser Ty				
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Lys Ser Tyr	Arg Ser His Lys Ly	s Tyr Tyr Lys	Lys Pro His His	: His	
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Trp Lys Asp	Gly Asn Cys Trp	Val Val Lys Ly:	s Lys Tyr Lys V	al Asp	
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		gag cat cat cct collu His His Pro Le 540			2107
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	585	590		595	
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		tc atc tgg tcg ct e lle Trp Ser Leu			2395
				4 , 4	

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ttt gcc gac cgc acg ccg cat cca gcg ctg acg gaa gca aaa cac cag Phe Ala Asp Arg Thr Pro His Pro Ala Leu Thr Glu Ala Lys His Gln cag cag ttt ttc cag ttc cgt tta tcc ggg caa acc atc gaa gtg acc Gin Gin Phe Phe Gin Phe Arg Leu Ser Gly Gin Thr lie Glu Val Thr age gaa tac etg tte egt eat age gat aac gag ete etg eac tgg atg Ser Glu Tyr Leu Phe Arg His Ser Asp Asn Glu Leu Leu His Trp Met gtg gcg ctg gat ggt aag ccg ctg gca agc ggt gaa gtg cct ctg gat Val Ala Leu Asp Gly Lys Pro Leu Ala Ser Gly Glu Val Pro Leu Asp gtc gct cca caa ggt aaa cag ttg att gaa ctg cct gaa cta ccg cag Val Ala Pro Gln Gly Lys Gln Leu Ile Glu Leu Pro Glu Leu Pro Gln ccg gag agc gcc ggg caa ctc tgg ctc aca gta cgc gta gtg caa ccg Pro Giu Ser Ala Gly Gin Leu Trp Leu Thr Val Arg Val Val Gin Pro . aac gcg acc gca tgg tca gaa gcc ggg cac atc agc gcc tgg cag cag Asn Ala Thr Ala Trp Ser Glu Ala Gly His Ile Ser Ala Trp Gin Gln tgg cgt ctg gcg gaa aac ctc agt gtg acg ctc ccc gcc gcg tcc cac Trp Arg Leu Ala Glu Asn Leu Ser Val Thr Leu Pro Ala Ala Ser His gcc atc ccg cat ctg acc acc agc gaa atg gat ttt tgc atc gag ctg Ala lle Pro His Leu Thr Thr Ser Glu Met Asp Phe Cys lle Glu Leu

ggt aat aag cgt tgg caa ttt aac cgc cag tca ggc ttt ctt tca cag Gly Asn Lys Arg Trp Gln Phe Asn Arg Gln Ser Gly Phe Leu Ser Gln atg tgg att ggc gat aaa aaa caa ctg ctg acg ccg ctg cgc gat cag Met Trp lie Gly Asp Lys Lys Gln Leu Leu Thr Pro Leu Arg Asp Gln ttc acc cgt gca ccg ctg gat aac gac att ggc gta agt gaa gcg acc Phe Thr Arg Ala Pro Leu Asp Asn Asp Ile Gly Val Ser Glu Ala Thr cgc att gac cct aac gcc tgg gtc gaa cgc tgg aag gcg gcg ggc cat Arg lle Asp Pro Asn Ala Trp Val Glu Arg Trp Lys Ala Ala Gly His tac cag gcc gaa gca gcg ttg ttg cag tgc acg gca gat aca ctt gct Tyr Gin Ala Giu Ala Ala Leu Leu Gin Cys Thr Ala Asp Thr Leu Ala gat gcg gtg ctg att acg acc gct cac gcg tgg cag cat cag ggg aaa Asp Ala Val Leu lle Thr Thr Ala His Ala Trp Gin His Gln Gly Lys acc tta ttt atc agc cgg aaa acc tac cgg att gat ggt agt ggt caa Thr Leu Phe lle Ser Arg Lys Thr Tyr Arg lle Asp Gly Ser Gly Gln atg gcg att acc gtt gat gtt gaa gtg gcg agc gat aca ccg cat ccg Met Ala lle Thr Val Asp Val Glu Val Ala Ser Asp Thr Pro His Pro gcg cgg att ggc ctg aac tgc cag ctg gcg cag gta gca gag cgg gta Ala Arg Ile Gly Leu Asn Cys Gln Leu Ala Gln Val Ala Glu Arg Val

	tta ggg ccg caa Leu Gly Pro Glr				3739
1080	10)85	1090		
	gac cgc tgg gat				3787
1095	Asp Arg Trp As 1100	p Leu Pro Lei	1105	iyi iili	
_	ccg agc gaa aad Pro Ser Glu Asn				3835
1110	1115		120	1125	
	cca cac cag tgg Pro His Gln Trp				3883
	1130	1135		1140	
	t caa cag caa ct Gin Gin Gin Leu		Ser His Arg His	s Leu	3931
· 114	5	1150	11:	55	
	a gaa ggc aca to Glu Gly Thr Trp				3979
1160	1	165	1170		
	gac gac tcc tgg Asp Asp Ser Trp 1180				4027
	c ggt cgc tac ca			•	4075
1190	Gly Arg Tyr His 1195		vai Trp Cys Gir 200	1:205	
taata ataa	accgggc aggcca	tgtc tgcccgta	tt togogtaagg	aaatccatta	4130
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Lys Pro His His Cys Asp Asp Tyr Lys Arg His Asp Asp Tyr Asp Ser Lys Lys Glu Tyr Trp Lys Asp Gly Asn Cys Trp Val Val Lys Lys Lys Tyr Lys Val Asp Arg Glu Asn Pro Gly Val Thr Gln Leu Asn Arg Leu Ala Ala His Pro Pro Phe Ala Ser Trp Arg Asn Ser Glu Glu Ala Arg Thr Asp Arg Pro Ser Gln Gln Leu Arg Ser Leu Asn Gly Glu Trp Arg Phe Ala Trp Phe Pro Ala Pro Glu Ala Val Pro Glu Ser Trp Leu Glu Cys Asp Leu Pro Glu Ala Asp Thr Val Val Val Pro Ser Asn Trp Gin Met His Gly Tyr Asp Ala Pro Ile Tyr Thr Asn Val Thr Tyr Pro lle Thr Val Asn Pro Pro Phe Val Pro Thr Glu Asn Pro Thr Gly Cys Tyr Ser Leu Thr Phe Asn Val Asp Glu Ser Trp Leu Gln Glu Gly Gln Thr Arg IIe IIe Phe Asp Gly Val Asn Ser Ala Phe His Leu Trp Cys Asn Gly Arg Trp Val Gly Tyr Gly Gln Asp Ser Arg Leu Pro Ser Glu

Phe Asp Leu Se	er Ala Phe Leu Ar	g Ala Gly Glu As	n Arg Leu Ala Val	
355	3	60	365	
Met Val Leu Arg 370	Trp Ser Asp Gly 375		Asp Gln Asp Met 380	t
Trp Arg Met Ser 385	r Gly lle Phe Arg 390	Asp Val Ser Leu 395	Leu His Lys Pro	400
Thr Thr Gln lie	Ser Asp Phe His 1	Val Ala Thr Arg F 410	Phe Asn Asp Asp 415	
Phe Ser Arg Ala	a Val Leu Glu Ala 20	Glu Val Gln Met 425	Cys Gly Glu Leu 430	
Arg Asp Tyr Le 435	u Arg Val Thr Val 4	Ser Leu Trp Gln 140	Gly Glu Thr Gln 445	
Val Ala Ser Gly 450	Thr Ala Pro Phe 455	Gly Gly Glu lle ll	e Asp Glu Arg 460	
Gly Gly Tyr Ala 465	Asp Arg Val Thr 470	Leu Arg Leu Asr 475		480
Lys Leu Trp Se	er Ala Glu lle Pro 485	Asn Leu Tyr Arg 490	Ala Val Val Glu 495	5
	a Asp Gly Thr Let	ı lle Glu Ala Glu 505	Ala Cys Asp Val 510	
Gly Phe Arg G	lu Val Arg lle Glu	Asn Gly Leu Leu 520	ı Leu Leu Asn Gly 525	,
Lys Pro Leu Le	eu lie Arg Gly Val 535	Asn Arg His Glu	His His Pro Leu 540	

His Gly Gln Val	Met Asp Glu G	In Thr Met Va	l Gin Asp ile L	.eu Leu
545	550		555	560
040				
Adat Lun Olm Ann	A A Dha Ann	Alo Vol Ara C	ve Sor His Tv	r Pro Asn
Met Lys Gln Asr			ys der ins iy	
	565	570		575
His Pro Leu Trp	Tyr Thr Leu C	ys Asp Arg Ty	r Gly Leu Tyr	Val Val
58	0	585		590
	. II. Ole The LE	a Ohi Mat Val	Dro Mot Asn	Ara lau
Asp Glu Ala Asr	i ile Giu i nr mi			
5 95		600	605	
Thr Asp Asp Pro	o Arg Trp Leu	Pro Ala Met S	er Glu Arg Val	Thr Arg
610	615		620	
0,0				
AA	A A A	Lio Dro Sor V	al lla lla Tra S	or Lou
Met Val Gln Arg		HIS PIO SEI V		
625	630		635	640
Gly Asn Glu Se	r Gly His Gly A	la Asn His As	p Ala Leu Tyr	Arg Trp
	645	650		655
	•			
U Luc Oca Val	A Dro Cor A	ra Dra Val Cla	Tyr Chi Chi C	elv Glv
lle Lys Ser Val			Tyl Glu Gly C	
66	30	665		670
Ala Asp Thr Th	r Ala Thr Asp I	le lle Cys Pro	Met Tyr Ala A	rg Val
675		680	685	5
A Oly A O	la Dra Dha Dra	. Ala Val Pra I	ve Tro Ser lle	lvelve
Asp Glu Asp G		_		Lyo Lyo
690	69	5	700	
			•	•
Trp Leu Ser Le	u Pro Gly Glu	Thr Arg Pro L	eu lle Leu Cys	Glu Tyr
705	710		715	720
A1- 11:- A1- A4-	+ Ohr Ann Son	an Gly Oly D	ha Ala Lua Tu	r Tro Clo
Ala His Ala Me				
	725	73	U	735

Ala Phe Arg Gin Tyr Pro Arg Leu Gin Gly Gly Phe Val Trp Asp Trp ·740 Val Asp Gln Ser Leu lle Lys Tyr Asp Glu Asn Gly Asn Pro Trp Ser Ala Tyr Gly Gly Asp Phe Gly Asp Thr Pro Asn Asp Arg Gln Phe Cys Met Asn Gly Leu Val Phe Ala Asp Arg Thr Pro His Pro Ala Leu Thr Glu Ala Lys His Gln Gln Gln Phe Phe Gln Phe Arg Leu Ser Gly Gln Thr lle Glu Val Thr Ser Glu Tyr Leu Phe Arg His Ser Asp Asn Glu Leu Leu His Trp Met Val Ala Leu Asp Gly Lys Pro Leu Ala Ser Gly Glu Val Pro Leu Asp Val Ala Pro Gln Gly Lys Gln Leu Ile Glu Leu Pro Glu Leu Pro Gln Pro Glu Ser Ala Gly Gln Leu Trp Leu Thr Val Arg Val Val Gln Pro Asn Ala Thr Ala Trp Ser Glu Ala Gly His Ile Ser Ala Trp Gln Gln Trp Arg Leu Ala Glu Asn Leu Ser Val Thr Leu Pro Ala Ala Ser His Ala Ile Pro His Leu Thr Thr Ser Glu Met Asp

Phe Cys Ile Glu Leu Gly Asn Lys Arg Trp Gln Phe Asn Arg Gln Ser Gly Phe Leu Ser Gln Met Trp Ile Gly Asp Lys Lys Gln Leu Leu Thr Pro Leu Arg Asp Gln Phe Thr Arg Ala Pro Leu Asp Asn Asp He Gly Val Ser Glu Ala Thr Arg lle Asp Pro Asn Ala Trp Val Glu Arg Trp Lys Ala Ala Gly His Tyr Gln Ala Glu Ala Ala Leu Leu Gln Cys Thr Ala Asp Thr Leu Ala Asp Ala Val Leu Ile Thr Thr Ala His Ala Trp Gin His Gin Gly Lys Thr Leu Phe lle Ser Arg Lys Thr Tyr Arg lle Asp Gly Ser Gly Gln Met Ala ile Thr Val Asp Val Glu Val Ala Ser Asp Thr Pro His Pro Ala Arg Ile Gly Leu Asn Cys Gln Leu Ala Gln Val Ala Glu Arg Val Asn Trp Leu Gly Leu Gly Pro Gln Glu Asn Tyr Pro Asp Arg Leu Thr Ala Ala Cys Phe Asp Arg Trp Asp Leu Pro Leu Ser Asp Met Tyr Thr Pro Tyr Val Phe Pro Ser Glu Asn Gly Leu Arg

Cys Gly Thr Arg Glu Leu Asn Tyr Gly Pro His Gln Trp Arg Gly Asp

1125

1130

1135

Phe Gln Phe Asn Ile Ser Arg Tyr Ser Gln Gln Gln Leu Met Glu Thr

1140

1145

1150

Ser His Arg His Leu Leu His Ala Glu Glu Gly Thr Trp Leu Asn lle

1155

1160

1165

Asp Gly Phe His Met Gly Ile Gly Gly Asp Asp Ser Trp Ser Pro Ser

1170

1175

1180

Val Ser Ala Glu Phe Gln Leu Ser Ala Gly Arg Tyr His Tyr Gln Leu

1185

1190

1195

1200

Val Trp Cys Gln Lys

1205

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<212> DNA

<213> Artificial Sequence

<220>

<223> cotG-linker 5' primer

<400> 27

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47

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<211> 49

<212> DNA

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WO 02/46388

PCT/KR01/02124

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catatato	ct ttttaattc acgcaagtct tttggatgaa caaacagctg ataaagcggt	300
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agatotoa	gc ggaacactta tacacttttt aaaaccgcgc gtactatgag ggtagtaagg	420
atcttcatc	c ttaacatatt tttaaaagga ggatttcaaa ttg ggc cac tat tcc Leu Gly His Tyr Ser	475
	1 5	
	ac atc gaa gaa gcg gtg aaa tcc gca aaa aaa gaa ggt tta	523
	Asp lie Glu Glu Ala Val Lys Ser Ala Lys Lys Glu Gly Leu	
	10 15 20	
aag gat	tat tta tac caa gag cct cat gga aaa aaa cgc agt cat aaa	571

Lys Asp Tyr Leu Tyr Gln Glu Pro His Gly Lys Lys Arg Ser His Lys aag tog cac ogc act cac aaa aaa tot ogc agc cat aaa aaa toa tac Lys Ser His Arg Thr His Lys Lys Ser Arg Ser His Lys Lys Ser Tyr tgc tct cac aaa aaa tct cgc agt cac aaa aaa tca ttc tgt tct cac Cys Ser His Lys Lys Ser Arg Ser His Lys Lys Ser Phe Cys Ser His aaa aaa tot ogo ago cac aaa aaa toa tac tgc tot cac aag aaa tot Lys Lys Ser Arg Ser His Lys Lys Ser Tyr Cys Ser His Lys Lys Ser cgc agc cac aaa aaa tcg tac cgt tct cac aaa aaa tct cgc agc tat Arg Ser His Lys Lys Ser Tyr Arg Ser His Lys Lys Ser Arg Ser Tyr aaa aaa tot tac ogt tot tac aaa aaa tot ogt ago tat aaa aaa tot Lys Lys Ser Tyr Arg Ser Tyr Lys Lys Ser Arg Ser Tyr Lys Lys Ser tgc cgt tct tac aaa aaa tct cgc agc tac aaa aag tct tac tgt tct Cys Arg Ser Tyr Lys Lys Ser Arg Ser Tyr Lys Lys Ser Tyr Cys Ser cac aag aaa aaa tot ogo ago tat aag aag toa tgo ogo aca cac aaa His Lys Lys Ser Arg Ser Tyr Lys Lys Ser Cys Arg Thr His Lys aaa tot tat ogt too cat aag aaa tac tac aaa aaa cog cac cac cac Lys Ser Tyr Arg Ser His Lys Lys Tyr Tyr Lys Lys Pro His His His tgc gac gac tac aaa aga cac gat gat tat gac agc aaa aaa gaa tac

Cys Asp Asp Ty	r Lys Arg His Asc 170	Asp Tyr Asp So 175	er Lys Lys Glu Ty 180	
	aat tgc tgg/gta Asn Cys Trp Val			it 1051
	cag gca tgc gct Gln Ala Cys Ala 20			ca 1099
-	ggc cag ctt ago Gly Gln Leu Ser 220	lle Lys Gly Thr G		1147
	a gcg gta cag ct Ala Val Gln Leu . 235			tg 1195 245
	gaa tat gtc aat a Glu Tyr Val Asn I 250			
	atc acc gtt ttc c y lle Thr Val Phe 5			1291
	gac aac ccg tcc sp Asn Pro Ser V 2			g 1339
	a aaa gag ctt gg Lys Glu Leu Gly 300			1387
atc tta aat gac	ggt aat cca aac	caa aat aaa gag	g aag gca aaa ga	aa 1435

lle Leu Asn Asp Gly Asn Pro Asn Gln Asn Lys Glu Lys Ala Lys Glu				
310	315	38	20	325
tto tto aag gaa	a atg tca agc ctt t	ac gga aac ac	g cca aac gtc att	1483
			Thr Pro Asn Val Ile	
The The Lyo	330	335	340	
	330	555	040	,
	•			4504
			ac tgg aag cgt gat	
Tyr Glu lle Ala	Asn Glu Pro Asn	Gly Asp Val As	sn Trp Lys Arg Asp	
3	45	350	355	
att aaa ccg ta	t gcg gaa gaa gtg	att too gtt ato	c cgc aaa aat gat	1579
lle Lys Pro Tyr	Ala Glu Glu Val II	le Ser Val Ile A	rg Lys Asn Asp	
360	5	365	370	
cca dac aac a	att atc att atc aga	acc out aca t	gg agc cag gat gtg	1627
				, , , , , , , , , , , , , , , , , , , ,
	e lie lie Val Gly Ti	in Giy iii iip	·	
375	380		385	
aat gat gct gc	c gat gac cag cta	a aaa gat gca	aac gtt atg gac gc	a 1675
Asn Asp Ala A	la Asp Asp Gln L	eu Lys Asp Ala	Asn Val Met Asp A	Ala
390	395	4	00	405
				-
ctt cat ttt tat g	jcc ggc aca cac	ggc caa ttt tta	cgg gat aaa gca	1723
Leu His Phe T	yr Ala Gly Thr His	Gly Gln Phe L	eu Arg Asp Lys Ala	1
	410	415	42	0
aac tat oca ct	c age aaa goa g	ca cct att tit d	tg aca gag tgg gga	1771
			al Thr Glu Trp Gly	•
` 4	125	430	435	
				1010
		•	ctt gat caa tog agg	
Thr Ser Asp Ala Ser Gly Asn Gly Gly Val Phe Leu Asp Gln Ser Arg				
440		445	450	
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Glu Trp Leu Ly 455		p Ser Lys Thi 460	r lle Ser Trp Va , 465	al Asn Trp		
aat ctt tct gat Asn Leu Ser A						1915
470	475		480		485	
	y Gly Trp Arg	Leu Ser Asp	t tta tot got toa Leu Ser Ala S	er-Gly Thr		1963
	490	•	495	500		
Phe Val Arg G			gat tog acg as Asp Ser Thr Ly		•	2011
Pro Glu Thr Pr		p Lys Pro Th	ica cag gaa aa r Gln Glu Asn (3ly lle Ser	t	2059
520		525	5	530		
			tg aac agc aac t Asn Ser Asn (it	2107
535		540	545			
			at acc acg gtt on the state of			2155
<u>5</u> 50	555		560		565	
gat_gtc_act_go Asp Val Thr A			aaa aac aaa Lys Asn Lys C 575			2203
att aac tat aa	ic tac aca ca	na ctt aga tac	c ggc_aat gtg a	nca tac aag		2251
			ys Gly Asn Val			
	585	590		595		
ttt gtg acg ttg	cat aaa cca	a aag caa ggi	t gca gat acc t	at ctg gaa		2299

Phe Val Thr Leu His Lys Pro Lys Gln Gly Ala Asp Thr Tyr Leu Glu 600 605 610 2347 ctt gga ttt aaa aac gga acg ctg gca ccg gga gca agc aca ggg aat Leu Gly Phe Lys Asn Gly Thr Leu Ala Pro Gly Ala Ser Thr Gly Asn 615 620 625 att cag ctt cgt ctt cac aat gat gac tgg agc aat tat gca caa agc 2395 lle Gin Leu Arg Leu His Asn Asp Asp Trp Ser Asn Tyr Ala Gin Ser 630 635 640 645 ggc gat tat tcc ttt ttc aaa tca aat acg ttt aaa aca acg aaa aaa 2443 Gly Asp Tyr Ser Phe Phe Lys Ser Asn Thr Phe Lys Thr Thr Lys Lys 650 660 655 atc aca tta tat gat caa gga aaa ctg att tgg gga aca gaa cca aat 2491 lle Thr Leu Tyr Asp Gln Gly Lys Leu lle Trp Gly Thr Glu Pro Asn 665 670 675

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<211> 677

<212> PRT

<213> Bacillus subtilis

<400> 35

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Lys Lys Glu Gly Leu Lys Asp Tyr Leu Tyr Gln Glu Pro His Gly Lys
20 25 30

Lys Arg Ser His Lys Lys Ser His Arg Thr His Lys Lys Ser Arg Ser
35 40 45

His Lys Lys Ser	Tyr Cys Ser His	s Lys Lys Ser	Arg Ser His Lys L	ys
50	. 55		60	
Ser Phe Cys Se	r His Lys Lys S	er Arg Ser His	s Lys Lys Ser Tyr (
65	70		75	80
Ser His Lys Lys	Ser Arg Ser Hi	s Lys Lys Ser	Tyr Arg Ser His L	
	85	. 90		95
Lys Ser Arg Ser	r Tyr Lys Lys Se	er Tyr Arg Ser	Tyr Lys Lys Ser A	rg
10	00	105	110	
Ser Tyr Lys Lys	Ser Cys Arg S	er Tyr Lys Lys	Ser Arg Ser Tyr l	_ys
115		120	125	
Lys Ser Tyr Cys	s Ser His Lys Ly	/s Lys Ser Arg	g Ser Tyr Lys Lys	Ser
130	135		140	
Cys Arg Thr Hi	s Lys Lys Ser T	yr Arg Ser His	s Lys Lys Tyr Tyr l	_ys
145	150		155	160
Lys Pro His His	s His Cys Asp /	Asp Tyr Lys A	rg His Asp Asp Ty	r Asp
	165	170		175
Ser Lys Lys Gl	u Tyr Trp Lys A	sp Gly Asn C	ys Trp Val Val Lys	Lys
1	80	185	190)
Lys Tyr Lys Gl	y Gly Gly Gly S	er Leu Gln Ala	a Cys Ala Ser Arg	Ser
195		200	205	
Gly Thr Lys Th	nr Pro Val Ala Ly	/s Asn Gly Gli	n Leu Ser IIe Lys (3ly
210	21	5	220	
Thr Gln Leu V	al Asn Arg Asp	Gly Lys Ala V	al Gln Leu Lys Gly	/ He
225	230		235	240

Ser Ser His Gly Leu Gln Trp Tyr Gly Glu Tyr Val Asn Lys Asp Ser Leu Lys Trp Leu Arg Asp Asp Trp Gly ile Thr Val Phe Arg Ala Ala Met Tyr Thr Ala Asp Gly Gly Ile Ile Asp Asn Pro Ser Val Lys Asn Lys Met Lys Glu Ala Val Glu Ala Ala Lys Glu Leu Gly Ile Tyr Val lle lle Asp Trp His lle Leu Asn Asp Gly Asn Pro Asn Gln Asn Lys Glu Lys Ala Lys Glu Phe Phe Lys Glu Met Ser Ser Leu Tyr Gly Asn Thr Pro Asn Val Ile Tyr Glu Ile Ala Asn Glu Pro Asn Gly Asp Val Asn Trp Lys Arg Asp Ile Lys Pro Tyr Ala Glu Glu Val Ile Ser Val lle Arg Lys Asn Asp Pro Asp Asn lle lle Val Gly Thr Gly Thr Trp Ser Gln Asp Val Asn Asp Ala Ala Asp Asp Gln Leu Lys Asp Ala Asn Val Met Asp Ala Leu His Phe Tyr Ala Gly Thr His Gly Gln Phe Leu Arg Asp Lys Ala Asn Tyr Ala Leu Ser Lys Gly Ala Pro Ile Phe

Val Thr Glu Trp Gly Thr Ser Asp Ala Ser Gly Asn Gly Gly Val Phe Leu Asp Gln Ser Arg Glu Trp Leu Lys Tyr Leu Asp Ser Lys Thr lle Ser Trp Val Asn Trp Asn Leu Ser Asp Lys Gln Glu Ser Ser Ser Ala Leu Lys Pro Gly Ala Ser Lys Thr Gly Gly Trp Arg Leu Ser Asp Leu Ser Ala Ser Gly Thr Phe Val Arg Glu Asn lie Leu Gly Thr Lys Asp Ser Thr Lys Asp lie Pro Glu Thr Pro Ala Lys Asp Lys Pro Thr Gln Glu Asn Gly lie Ser Val Gln Tyr Arg Ala Gly Asp Gly Ser Met Asn Ser Asn Gln lle Arg Pro Gln Leu Gln lle Lys Asn Asn Gly Asn Thr Thr Val Asp Leu Lys Asp Val Thr Ala Arg Tyr Trp Tyr Asn Ala Lys Asn Lys Giy Gin Asn Val Asp Cys Asp Tyr Ala Gin Leu Gly Cys Gly Asn Val Thr Tyr Lys Phe Val Thr Leu His Lys Pro Lys Gln Gly Ala Asp Thr Tyr Leu Glu Leu Gly Phe Lys Asn Gly Thr Leu Ala Pro Gly

Ala Ser Thr Gly Asn lle Gln Leu Arg Leu His Asn Asp Asp Trp Ser

625

635

640

Asn Tyr Ala Gln Ser Gly Asp Tyr Ser Phe Phe Lys Ser Asn Thr Phe

645

630

650

655

Lys Thr Thr Lys Lys lie Thr Leu Tyr Asp Gin Gly Lys Leu lie Trp

660

665

670

Gly Thr Glu Pro Asn

675

<210> 36

<211> 24

<212> PRT

<213> Hepatitis B virus

<400> 36

Met Gin Trp Asn Ser Thr Thr Phe His Leu Gin Asp Pro Arg Val Arg

1

5

10

15

Gly Leu Tyr Phe Pro Ala Gly Gly

20

<210> 37

<211> 30

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<213> Artificial Sequence

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<400> 42

cgaagcttga attcttattt gtagagctca tc 32

INTERNATIONAL SEARCH REPORT

International application No. PCT/KR01/02124

A. CLA	ASSIFICATION OF SUBJECT MATTER			
	C7 C12N 15/00			
According to	International Patent Classification (IPC) or to both na	tional classification and IPC		
	LDS SEARCHED		- M M	
	ocumentation searched (classification system followed	by classification symbols)		
IPC(7) C12				
	on searched other than minimum documentation to the	extent that such documents are included in the	fields searched	
	ent and application for invention since 1975			
Electronic da Kipass, Me	ta base consulted during the intertnational search (nam dline, Delphion, PAJ	e of data base and, where practicable, search ter	ms used)	
C. DOCU	MENTS CONSIDERED TO BE RELEVANT			
Category*	Citation of document, with indication, where ap	opropriate, of the relevant passages	Relevant to claim No.	
Y	US 5,766,914(Michigan State University), 16 Jun 1	998 (see the whole document)	1-4,6,10-14,17-26 28, 30-34, 40-42	
Y	Hiroshi Ichikawa et al., "Combined action of transcription factors regulates genes encoding spore coat proteins of Bacillus subtilis", J Bioll Chem., May 2000 275(18):13849-55 (see the whole document)			
Α	US 5,837,500(Dyax, Corp.) 17 Nov. 1998 (see the	whole document)	7	
Further	r documents are listed in the continuation of Box C.	X See patent family annex.		
'A" document	categories of cited documents: It defining the general state of the art which is not considered articular relevence	"T" later document published after the internati date and not in conflict with the applicati	onal filing date or priority	
E" earlier ap	ention ned invention cannot be			
"L" document which may throw doubts on priority claim(s) or which is step when the document is taken alone				
special reason (as specified) "O" document referring to an oral disclosure, use, exhibition or other "Y" document of particular relevence; the claimed invention considered to involve an inventive step when the decomposition of the combined with one or more other such documents, such				
	t published prior to the international filing date but later priority date claimed	being obvious to a person skilled in the art "&" document member of the same patent famil		
Date of the act	rual completion of the international search	Date of mailing of the international search rep	ort	
	0 APRIL 2002 (30.04.2002)	30 APRIL 2002 (30.04.2002)		
Korean Intelle	iling address of the ISA/KR ectual Property Office	Authorized officer	As man	
Government	Complex-Daejeon, 920 Dunsan-dong, Seo-gu,	AHN, Mi Jung	A la sezza A	
Dacjeon Meti	ropolitan City 302-701, Republic of Korea 82-42-472-7140	Autia' ian lank		

INTERNATIONAL SEARCH REPORT

Information on patent family members

International application No.

PCT/KR01/02124

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